SEARCH REQUEST FORM

469

Scientific and Technical Inf rmation Center

Requester's Full Name: PATE Art Unit: 1 (24 Phone	LSUDTIAKE Number 30 8 47	RExaminer #: 7)	18 Date: 5/16/02
Mail Box and Bldg/Room Location	on CIMI AFIT Po	Sults Format Professed	6 046 526
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If more than one search is sub	mitted, please priorit	ize searches in order o	f need.
Please provide a detailed statement of the	e search topic, and describ	e as specifically as possible the	subject matter to be searched
the district species of structures.	KCVWUIUS, SVIRONVING ACTA	invinc and redictor numbers of	and an article of the N
utility of the invention. Define any term known. Please attach a copy of the cover	succi, pertinent claims, ar	id abstract.	
SUBSTITUTED AR	YLAMINE?	DERIVATIVES &	METHEN OF USE
Inventors (please provide full names):	GUOQIA	IG CHEN	
(Produce provide rail names).	<u> </u>	GCHEN	<u> ८ दूर</u>
			<u> </u>
Earliest Priority Filing Date:		cr	-C112-C112-C-
For Sequence Searches Only Please inclu appropriate serial number.	ide all pertinent information	(parent, child, divisional, or issu	ed patent numbers) along with the
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STAFF USE ONLY	Type of Search	Vendors and cost	where applicable
Searcher: Selevy 6.477	NA Sequence (#)	STN !	
Searcher Phone #:	AA Sequence (#)	Dialog	• • • • • • • • • • • • • • • • • • • •
Searcher Location:	Structure (#)	Questel/Orbit	
Date Searcher Picked Up:	Bibliographic	Dr.Link	·
Date Completed: 05-31-02	Litigation	Lexis/Nexis	
Searcher Prep & Review Time:	Fulltext	Sequence Systems	·
Clerical Prep Time:	Patent Family	WWW/Internet	
Online Time:	Other	Other (specify)	
PTO-1590 (8-01)			\$

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REGISTRY - ENTERED AT 14:37:32 ON 31 MAY 2002)
L4
                STR
           10
               G6
           N~G5~Cb
           11 12 13
VAR G1=C/N
VAR G2=C/N/O/S
REP G3=(0-1) C
VAR G4=O/S
REP G5=(0-1) CH2
VAR G6=H/ET/ME/I-PR/N-PR/I-BU/N-BU/S-BU/T-BU/CY
NODE ATTRIBUTES:
                  \mathbf{AT}
                       9
NSPEC
       IS RC
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
GRAPH ATTRIBUTES:
RSPEC I
NUMBER OF NODES IS 14
STEREO ATTRIBUTES: NONE
L5
                STR
               14
                                                     Str. 2
           N~G5~Cb
           11
             12 13
VAR G1=C/N
VAR G2=C/N/O/S
REP G3=(0-1) N
VAR G4=O/S
REP G5=(0-1) CH2
VAR G6=H/ET/ME/I-PR/N-PR/I-BU/N-BU/S-BU/T-BU/CY
NODE ATTRIBUTES:
NSPEC
       IS RC
                  AΤ
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
GRAPH ATTRIBUTES:
RSPEC I
NUMBER OF NODES IS 14
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2544 SEA FILE=REGISTRY SSS FUL L4 OR L5 & Temp Sources 7 days
STEREO ATTRIBUTES: NONE
L7
L8
                STR
           10
              14
              G6
              ~ N~~ C
           N-√G5-√Cb
          11 12
VAR G1=C/N
VAR G2=C/N/O/S
REP G3=(0-1) C
VAR G4=O/S
REP G5=(0-1) CH2
VAR G6=H/ET/ME/I-PR/N-PR/I-BU/N-BU/S-BU/T-BU/CY
NODE ATTRIBUTES:
NSPEC
       IS RC
                 ΑT
DEFAULT MLEVEL IS ATOM
GGCAT
       IS UNS AT 13
DEFAULT ECLEVEL IS LIMITED
GRAPH ATTRIBUTES:
RSPEC I
NUMBER OF NODES IS 14
STEREO ATTRIBUTES: NONE
L9
           1687 SEA FILE=REGISTRY SUB=L7 SSS FUL L8
L10
               STR
           10
              14
              G6
           N~G5~Cb
          11 12
VAR G1=C/N
VAR G2=C/N/O/S
REP G3=(0-1) N
VAR G4=O/S
REP G5=(0-1) CH2
VAR G6=H/ET/ME/I-PR/N-PR/I-BU/N-BU/S-BU/T-BU/CY
NODE ATTRIBUTES:
NSPEC
       IS RC
                 AΤ
DEFAULT MLEVEL IS ATOM
GGCAT
       IS UNS AT 13
DEFAULT ECLEVEL IS LIMITED
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GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

616 SEA FILE=REGISTRY SUB=L7 SSS FUL L10 L11

L12 2132 SEA FILE=REGISTRY ABB=ON PLU=ON L9 OR L11

35616002 1/NC (1) 2009 L12 AND 1/NC 2

(FILE CAPLOS ENTERED AT 14:41:28 ON 31 MAY 2002)

392 SEA ABB=ON PLU=ON L13 OR L13/D L14

L15 170 SEA ABB=ON PLU=ON L14 AND (PROPHYLACT? OR PROPHYLAX?

OR TREAT? OR THERAP?)

43 SEA ABB=ON PLU=ON L15 AND (DISEAS? OR DISORDER OR L16 MALAD?)

L16 ANSWER 1 OF 43 CAPLUS COPYRIGHT 2002 ACS

2002:275966 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 136:294739

Preparation of pyridinyl-substituted benzamides TITLE:

as Apo B secretion inhibitors

INVENTOR(S): Takasugi, Hisashi; Terasawa, Takeshi; Inoue,

Yoshikazu; Nakamura, Hideko; Nagayoshi, Akira; Ohtake, Hiroaki; Furukawa, Yoshiro; Mikami, Masafumi; Hinoue, Kazumasa; Ohtsubo, Makoto

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co., Ltd., Japan; Daiso

Co., Ltd.

SOURCE: PCT Int. Appl., 266 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA		KIND DATE				APPLICATION NO.						DATE					
									_								
WO	2002	0288	35	A1 20020411					W	0 20	01-J	1	20010928				
	W:	ΑE,	AG,	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	ΗU,	ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	KR,	ΚZ,	LC,	
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NO,	
		NZ,	PH,	PL,	PT,	RO,	RU,	SD									
	RW:							•		•				AT,			
		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	
		TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	G₩,	ML,	MR,	ΝĒ,	SN,	
		TD,	TG														
PRIORITY	APP	LN.	INFO	.:				1	AU 2	000-	583		Α	2000	1005		
								7	AU 2	001-	6666		Α	2001	0727		

OTHER SOURCE(S): MARPAT 136:294739

GΙ

$$\begin{array}{c|c}
R^{1} \\
 & \downarrow \\
 & \downarrow$$

Title compds. I [wherein R1 and R2 = independently alkyl, alkenyl, AB acyl, amino, (cyclo)alkoxy, aryl(oxy), sulfoxy, mercapto, sulfo, H, halo, NO2, CN, or OH; or R1R2 = a ring; Q1 = N or CH; L = (un) substituted unsatd. 3 to 10-membered heterocyclic group; X =(un) substituted monocyclic (hetero) arylene; Y = (A1)m(A2)n(A4)k; Z =direct bond, CH2, NH, or O; R = H or alkyl; A1 = (un)substituted alkylene or alkenylene; A2 = NR3, CONR3, NHCONH, CO2, O, O(CH2)2NR3, S, SO, or SO2; A4 = alkylene, alkenylene, or alkynylene; R3 = H or suitable substituent; k, m, and m = independently 0 or 1; or a salt thereof] were prepd. as apolipoprotein B (Apo B) secretion inhibitors. For example, to a suspension of N-(4-aminophenyl)-4'-(trifluoromethyl)-1,1'-biphenyl-2-carboxamide, 2-pyridinylacetic acid.bul.HCl, and HOBT.bul.H2O in CH2Cl2 was added to WSC.bul.HCl, followed by TEA at 5.degree.C. The mixt. was stirred at room temp. for 24 h and worked up to give II. The latter inhibited Apo B secretion by 100% at 10-6 M in HepG2 cells and lowered cholesterol by 83% and triglyceride by 35% after 2 h at a dose of 32 mg/kg in ddY-mice. I are useful for the prophylaxis and treatment of diseases or conditions resulting from elevated circulating levels of Apo B, such as hyperlipemia, hyperlipidemia, hyperlipoproteinemia, hypoalphalipoproteinemia, hypercholesterolemia, hypertriglyceridemia, atherosclerosis, pancreatitis, non-insulin dependent diabetes mellitus, obesity, coronary heart diseases, myocardial infarction, stroke, restenosis, and Syndrome X.

II

(Preparation); RACT (Reactant or reagent); USES (Uses) (Apo B inhibitor; prepn. of pyridinyl-substituted benzamides as Apo B secretion inhibitors for treatment of obesity, NIDDM, and related conditions) 408364-90-9P, N-[4-[[2-(2-Pyridinyl)ethyl]amino]phenyl]-2-[3-(trifluoromethyl)anilino]benzamide 408365-53-7P, N-[4-[[2-[3-(Trifluoromethyl)anilino]benzoyl]amino]benzyl]-2pyridinecarboxamide 408365-66-2P, N-[4-[Methyl[2-(2pyridinyl)ethyl]amino]phenyl]-2-[3-(trifluoromethyl)anilino]benzamid e 408369-40-4P, N-[3-Fluoro-4-[[2-(2pyridinyl)ethyl]amino]phenyl]-2-[3-(trifluoromethyl)anilino]benzamid RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Apo B inhibitor; prepn. of pyridinyl-substituted benzamides as Apo B secretion inhibitors for treatment of obesity, NIDDM, and related conditions) THERE ARE 5 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT: 5 THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L16 ANSWER 2 OF 43 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2002:237356 CAPLUS 136:263090 DOCUMENT NUMBER: TITLE: Preparation of cyclic amine derivatives for inhibition of the action of chemokines such as MIP-1.alpha. and/or MCP-1 on target cells Shiota, Tatsuki; Kataoka, Ken-Ichiro; Imai, INVENTOR(S): Minoru; Tsutsumi, Takaharu; Sudoh, Masaki; Sogawa, Ryo; Morita, Takuya; Hada, Takahiko; Muroga, Yumiko; Takenouchi, Osami; Furuya, Minoru; Endo, Noriaki; Tarby, Christine M.; Moree, Wilna; Teig, Steven Teijin Limited, Japan; Dupont Pharmaceuticals PATENT ASSIGNEE(S): Research Laboratories SOURCE: U.S., 364 pp., Cont. of U.S. Ser. No. 554,562. CODEN: USXXAM DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: DATE PATENT NO. KIND DATE APPLICATION NO. _____ -----____ US 2001-905078 20010716 US 6362177 B1 20020326 PRIORITY APPLN. INFO.: US 2000-554562 A3 20000516 OTHER SOURCE(S): MARPAT 136:263090 GΙ

TT

308-4994 Shears Searcher :

$$\begin{array}{c|c}
R^{1} & O & R^{4} \\
 & CH_{2} \\
 & M & CH_{2} \\
 & M & R^{3}
\end{array}$$

$$\begin{array}{c|c}
CH_{2} \\
 & P \\
 & R^{5}
\end{array}$$

$$\begin{array}{c|c}
CH_{2} \\
 & Q \\
 & Q$$

$$\begin{array}{c|c} C1 \\ & H & O \\ & \downarrow & \downarrow \\ & N & \downarrow \\$$

IT

The title compds. [I; R1 = (un)substituted Ph, cycloalkyl, AΒ heteroaryl, etc.; R2 = H, alkyl, alkoxycarbonyl, etc.; j = 0-2; k =0-2; m = 3-4 and k+m = 5 or 6; n = 0-1; R3 = H, alkyl; R4, R5 = H, OH, Ph, etc.; p, q = 0-1; G = CO, SO, CO2, etc.; R6 = Ph, cycloalkyl, cycloalkenyl, etc.] and their pharmaceutically acceptable acid addn. salts which inhibit the action of chemokines such as MIP-1.alpha. and/or MCP-1 on target cells and may be useful as a therapeutic drug and/or preventative drug in diseases, such as atherosclerosis, rheumatoid arthritis, and the like where blood monocytes and lymphocytes infiltrate into tissues, were prepd. Thus, reaction of N-benzoylglycine with 3-amino-1-(4-chlorobenzyl)pyrrolidine.2HCl in the presence of 3-ethyl-1-[3-(dimethylaminopropyl)]carbodiimide. HCl, 1-hydroxybenzotriazole and Et3N in CHCl3 afforded 95% II which showed 50-80% inhibition of MIP-1.alpha. binding to THP-1 cells at

10 .mu.M. 226241-50-5P, Benzamide, 5-chloro-2-[[(4ethoxyphenyl)methyl]amino]-N-[2-[[[1-[(4-ethoxyphenyl)methyl]-4piperidinyl]methyl]amino]-2-oxoethyl]- 226241-52-7P, Benzamide, 5-bromo-2-[[(4-ethoxyphenyl)methyl]amino]-N-[2-[[[1-[(4ethoxyphenyl)methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]-226241-63-0P, Benzamide, 5-chloro-2-[[(4ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]-4piperidinyl]methyl]amino]-2-oxoethyl]- 226241-64-1P, Benzamide, 5-chloro-2-[[[4-(1-methylethyl)phenyl]methyl]amino]-N-[2-[[[1-[[4-(1-methylethyl)phenyl]methyl]-4-piperidinyl]methyl]amino]-2oxoethyl] - 226241-65-2P, Benzamide, 5-chloro-N-[2-oxo-2-[[[1-[(4-propoxyphenyl)methyl]-4-piperidinyl]methyl]amino]ethyl]-2-[[(4-propoxyphenyl)methyl]amino]- 226241-66-3P, Benzamide, 5-bromo-2-[[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[[1-[(4-ethylphenyl)methyl]amino]-N-[2-[[1-[(4-[(4-ethylphenyl)methyl]amino]-N-[2-[[1-[(4-[(4-ethylphenyl)methyl]amino]-N-[2-[[1-[(4-[(4-[thylphenyl)methyl]amino]-N-[2-[[1-[(4-[thylphenyl]methyl]amino]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenyl]methyl]methyl]-N-[2-[[1-[(4-[thylphenethylphenyl)methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]-226241-67-4P, Benzamide, 5-bromo-2-[[[4-(1methylethyl) phenyl] methyl] amino] -N-[2-[[[1-[4-(1methylethyl)phenyl]methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]-226241-68-5P, Benzamide, 5-bromo-N-[2-oxo-2-[[[1-[(4propoxyphenyl)methyl]-4-piperidinyl]methyl]amino]ethyl]-2-[[(4propoxyphenyl)methyl]amino]- 226241-69-6P, Benzamide, 5-bromo-2-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]amino]]-N-[2-[[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[4-(methylthio)phenyl]methyl]methyl]-N-[2-[[4-(methylthio)phenyl]methyl]-N-[2-[[4-(methylthio)phenyl]methyl]-N-[2-[[4-(methylthio)phenyl]methyl]-N-[4-(methylthio)phenyl-N-[4-(methylthio)phenyl-N-[4-(methylthio)phenyl-N-[4-(methylthio)phenyl-N-[4-(methylthio)phenyl-N-[4-(methylthio)phenyl-N-[4-(methylthio)phenyl-N-[4-(methylthio)phenyl-N-[4-(methylth

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(methylthio)phenyl]methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]-
     226241-82-3P, Benzamide, 5-chloro-2-[[(4-hydroxy-3-
     methoxyphenyl)methyl]amino]-N-[2-[[[1-[(4-hydroxy-3-
     methoxyphenyl)methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]-
     226241-83-4P, Benzamide, 5-bromo-2-[[(4-hydroxy-3-
     methoxyphenyl)methyl]amino]-N-[2-[[[1-[(4-hydroxy-3-
     methoxyphenyl)methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]-
     226242-54-2P, Benzamide, 5-chloro-2-[[[4-
     (methylthio) phenyl] methyl] amino] -N-[2-[[[1-[[4-
     (methylthio)phenyl]methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]-
     226243-23-8P, Benzamide, 5-bromo-2-[[(4-
     butylphenyl)methyl]amino]-N-[2-[[[1-[(4-butylphenyl)methyl]-4-
     piperidinyl]methyl]amino]-2-oxoethyl]- 226243-25-0P,
     Benzamide, 5-bromo-N-[2-oxo-2-[[[1-[(4-propylphenyl)methyl]-4-
     piperidinyl]methyl]amino]ethyl]-2-[[(4-propylphenyl)methyl]amino]-
     226243-27-2P, Benzamide, 2-[[(4-butylphenyl)methyl]amino]-N-
     [2-[[1-[(4-butylphenyl)methyl]-4-piperidinyl]methyl]amino]-2-
     oxoethyl]-5-chloro- 226243-29-4P, Benzamide,
     5-chloro-N-[2-oxo-2-[[[1-[(4-propylphenyl)methyl]-4-
     piperidinyl]methyl]amino]ethyl]-2-[[(4-propylphenyl)methyl]amino]-
     226245-19-8P, Benzamide, 5-bromo-2-[[(4-
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     RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (prepn. of cyclic amine derivs. for inhibition of action of
        chemokines such as MIP-1.alpha. and/or MCP-1 on target cells)
REFERENCE COUNT:
                          24
                                THERE ARE 24 CITED REFERENCES AVAILABLE
                                FOR THIS RECORD. ALL CITATIONS AVAILABLE
                                IN THE RE FORMAT
L16 ANSWER 3 OF 43 CAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                          2002:157743 CAPLUS
DOCUMENT NUMBER:
                          136:217047
                          Preparation of novel phenylalanine derivatives
TITLE:
                         having .alpha.4 integrin-inhibitory activity
                         Makino, Shingo; Okuzumi, Tatsuya; Yoshimura,
INVENTOR(S):
                          Toshihiko; Satake, Yuko; Suzuki, Nobuyasu;
                          Izawa, Hiroyuki; Sagi, Kazuyuki; Chiba, Akira;
                          Nakanishi, Eiji; Murata, Masahiro; Tsuji,
                          Takashi
                          Ajinomoto Co., Inc., Japan
PATENT ASSIGNEE(S):
                          PCT Int. Appl., 137 pp.
SOURCE:
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
                          Japanese
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND DATE
                                            APPLICATION NO.
                                                              DATE
                             _____
                                            -----
                      ____
                                                             20010815
                                            WO 2001-JP7039
     WO 2002016329
                             20020228
                      A1
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,
             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ,
             NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR,
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TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ,

MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN,

TD, TG

PRIORITY APPLN. INFO.:

JP 2000-248728 A 20000818

JP 2001-147451 A 20010517

OTHER SOURCE(S):

MARPAT 136:217047

GI

$$\begin{array}{c} J \\ A \\ J' \\ D-T-N \\ CH \\ CO-B \\ C \end{array}$$

AB Phenylalanine derivs. [I; A = Q, Q1, Q2, Q3; wherein Arm = cyclic alkyl or arom. ring contg. 1-4 heteroatom(s) selected from O, S, and N; U, V, X = CO, SO2, CR5R6, C(:CR5R6), C:S, S:O, P(O)OH, P(O)H; W = CR7, N; wherein R1 - R7 = H, H, halo, OH, (un) substituted lower alkyl, alkenyl, or alkynyl, cycloalkyl optionally contg. a heteroatom in the ring, aryl, heteroaryl, etc.; B = HO, lower alkoxy, hydroxyamino; C = H, lower alkyl, alkenyl, alkynyl, cycloalkyl-lower alkyl (optionally contg. an heteroatom in the ring), aryl-lower alkyl, heteroaryl-lower alkyl; D = lower alkyl, alkenyl, alkynyl, cycloalkyl or cycloalkyl-lower alkyl (optionally contg. an heteroatom in the ring), aryl, aryl-lower alkyl, heteroaryl-lower alkyl, lower alkoxy, cycloalkyl-lower alkoxy (optionally contg. a heteroatom in the ring), aryloxy, heteroaryloxy, etc.; or C and D are linked to each other to form a ring optionally contg. 1 or 2 O, N, or S atom(s); T = CO, C:S, SO, SO2, NHCO, NHCS; J, J' = H, halo, lower alkyl, lower alkoxy, NO2] are prepd. by the solid phase method using Wang resin. These compds. are useful for the treatment or prevention of inflammatory disease states related to the .alpha.4 integrin-dependent adhesion process, e.g. rheumatoid arthritis, inflammatory bowel disease, systemic lupus erythematosus, multiple sclerosis, Sjoegren's syndrome, asthma, psoriasis, allergy, diabetes, cardiovascular diseases, atherosclerosis, restenosis, tumor proliferation, tumor metastasis, and transplant rejection. Thus, a soln. of Fmoc-Phe(4-NO2)-OH, 2,6-dichlorobenzoyl chloride, and pyridine in N-methylpyrrolidone was added to Wang resin and stirred at room temp. for 16 h to give Fmoc-Phe(4-NO2)-Wang resin which was deprotected by 20% piperidine in DMF at room temp. for 15 min to afford H-Phe(4-NO2)-Wang resin and then acylated by 2,6-dichlorobenzoyl chloride and 2,6-lutidine in N-methylpyrrolidone at room temp. for 16 h to give 2,6-dichlorobenzoyl-Phe(4-NO2)-Wang resin. The latter compd.-bound resin was reduced by SnCl2.2H2O in EtOH/N-methylpyrrolidone at room temp. for 16 h to 2,6-dichlorobenzoyl-Phe(4-NH2)-Wang resin which

```
was cyclocondensed with Me 2-isocyanatobenzoate in
     N-methylpyrrolidone at room temp. for 16 h to give
     2,6-dichlorobenzoyl-Phe(4-Q)-Wang resin (Q = 1,2,3,4-tetrahydro
     quinazolin-3-yl) and then methylated by Me iodide in the presence of
     18-crown-6 ether and K2CO3 in N-methylpyrrolidone at room temp. for
     3 days to give 2,6-dichlorobenzoyl-Phe(4-Q)-Wang resin (Q =
     1-methyl-1,2,3,4-tetrahydroquinazolin-3-yl). Resin-cleavage
     reaction with 5% aq. CF3CO2H at room temp. for 1 h gave
     2,6-dichlorobenzoyl-Phe(4-Q)-OH(Q = 1-methyl-1,2,3,4-
     tetrahydroquinazolin-3-yl) (II). II and 2-chloro-6-methylbenzoyl-
     Phe (4-Q) -OH (Q = 1-methyl-1,2,3,4-tetrahydroquinazolin-3-yl)
     inhibited the binding of human recombinant VCAM-1 to human T cell
     Jurikat (ATCC TIB-152) cell expressing integrin .alpha.4.beta.1 with
     IC50 of 1.0 and 0.2 nM, resp.
     401905-99-5DP, Wang resin-bound
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent)
        (rejection of)
                               THERE ARE 5 CITED REFERENCES AVAILABLE FOR
REFERENCE COUNT:
                               THIS RECORD. ALL CITATIONS AVAILABLE IN
                               THE RE FORMAT
L16 ANSWER 4 OF 43 CAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                         2001:923748 CAPLUS
DOCUMENT NUMBER:
                         136:53544
                         .beta.-amino acid nitrile derivs. useful for the
TITLE:
                         treatment of diseases which
                         are assocd. with cysteine proteases
                         Gabriel, Tobias; Pech, Michael; Rodriguez
INVENTOR(S):
                         Sarmiento, Rosa Maria
                         F. Hoffmann-La Roche A.-G., Switz.
PATENT ASSIGNEE(S):
SOURCE:
                         PCT Int. Appl., 91 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                          APPLICATION NO.
                                                            DATE
     PATENT NO.
                     KIND
                           DATE
                                          -----
                      ____
                           -----
                                         WO 2001-EP6541 20010608
                     A1
                            20011220
     WO 2001096285
            AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CO,
             CU, CZ, DE, DK, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU,
             ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
            LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU,
             ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD,
             TG
                            20020207
                                           US 2001-872927
                                                            20010601
     US 2002016361
                      Α1
                                        EP 2000-112577 A 20000614
```

308-4994 Searcher : Shears

MARPAT 136:53544

PRIORITY APPLN. INFO.:

OTHER SOURCE(S):

GI

$$R^{2}$$
 O R^{4} CN R^{1} CN R^{3} C R^{3} C R^{3}

Compds. of formula I [R1 = H, aryl, C(0)Ra, or SO2Rb (Ra = lower AB alkyl, lower-alkoxy, cycloalkyl, cycloalkyl-lower-alkyl, cycloalkyl-lower alkoxy, cycloalkoxy, aryl, aryloxy, etc.; Rb = aryl, aryl-lower-alkyl, or heteroaryl); R2, R3, R4 = H or lower-alkyl; R5 = H, lower-alkyl, cycloalkyl, or aryl; n = 1,2] were prepd.. Thus, $(1R, 2R) - (2-\{(S) - [cyano(3$ hydroxyphenyl)methyl]carbamoyl}cyclohexyl)carbamic acid benzyl ester (II) was produced from (1R, 2R)-2-benzyloxycarbonylaminocyclohexane carboxylic acid and (S)-2-amino-2-(3-hydroxyphenyl)acetonitrile. was assayed against cathepsins K, S, L, and B and the inhibitory activity (IC50) was detd. to be 0.005, >10, 4.7, and 4.6 .mu.Mol/L, resp. The compds. and pharmaceutically acceptable salts and/or pharmaceutically acceptable esters thereof are useful for the treatment of diseases which are assocd. with cysteine proteases such as osteoporosis, osteoarthritis, rheumatoid arthritis, tumor metastasis, glomerulonephritis, atherosclerosis, myocardial infarction, angina pectoris, instable angina pectoris, stroke, plaque rupture, transient ischemic attacks, amaurosis fugax, peripheral arterial occlusive disease, restenosis after angioplasty and stent placement, abdominal aortic aneurysm formation, inflammation, autoimmune disease, malaria, ocular fundus tissue cytopathy and respiratory disease. A discussion of pharmaceutical compns. is also included.

II

IT 381240-25-1P 381240-26-2P 381240-27-3P 381240-28-4P 381240-30-8P 381240-32-0P 381240-33-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of beta-amino acid nitrile derivs. useful for the treatment of diseases which are assocd. with

cysteine proteases)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR

THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L16 ANSWER 5 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:816647 CAPLUS

DOCUMENT NUMBER:

135:357948

TITLE:

Preparation of heterocyclic compounds as

phosphodiesterase V (PDE V) inhibitors

INVENTOR(S):

Yamada, Koichiro; Matsuki, Kenji; Omori, Kenji; Kikkawa, Kohei

PATENT ASSIGNEE(S):

Tanabe Seiyaku Co., Ltd., Japan

SOURCE:

PCT Int. Appl., 207 pp.

DOCUMENT TYPE:

Patent

CODEN: PIXXD2

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.				KIND DATE					A.	PPLI	CATI	ο.	DATE			
			-													
WO	2001	0834	60	Α	1	2001	1108		M	0 20	01-J	P203	4	2001	0315	
	W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,
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		GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KR,	ΚZ,	LC,	LK,
		LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	ΜW,	MX,	MZ,	NO,	NZ,
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,
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		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,
		TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,
		TG														
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PRIORITY APPLN. INFO.:

JP 2000-130371 A 20000428

OTHER SOURCE(S):

MARPAT 135:357948

GT

$$x$$
 R^2
 COR^3

AB Compds. of the general formula (I) or pharmacol. acceptable salts thereof [wherein X is :CH or N; Y is NH, NR4, S, O, CH:N, N:CH, N:N, CH:CH, or the like; R1 is lower alkoxy, amino, a nitrogenous heterocyclic group, or a hydroxyl group substituted with a heterocyclic group (wherein each group may be substituted); R2 is either a lower alkylamino or lower alkoxy group which may be substituted with aryl, or a lower alkoxy group substituted with a nitrogenous arom. heterocyclic group; and R3 is aryl, a nitrogenous heterocyclic group, lower alkyl, lower alkoxy, lower cycloalkoxy, a hydroxyl group substituted with a nitrogenous heterocyclic group, or amino (wherein each group may be substituted), or alternatively, R3

and the substituent of Y may be united to form a lactone ring] or pharmacol. acceptable salts thereof are prepd. These compds. exhibit excellent PDE V inhibitory activity and are useful as preventive or therapeutic agents for various. diseases due to dysfunction of the signal transduction through cGMP, in particular impotence, pulmonary hypertension, and diabetic renal failure paralysis (no data). Thus, 2-(hydroxymethyl)pyridine was treated wit NaH in THF at room temp. for 30 min and then condensed with 2-chloro-5-(3,4,5trimethoxyphenylcarbonyl)-4-(3-chloro-4methoxybenzylamino)pyrimidine (prepn. given) in THF at room temp. for 1 h to give 2-(2-pyridylmethoxy)-5-(3,4,5trimethoxyphenylcarbonyl)-4-(3-chloro-4methoxybenzylamino)pyrimidine. 330784-43-5P 372115-79-2P 372115-80-5P 372115-84-9P 372115-85-0P 372115-86-1P 372115-93-0P 372115-94-1P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (prepn. of heterocyclic compds. as phosphodiesterase V inhibitors preventive or therapeutic agents for various diseases due to dysfunction of signal transduction through cGMP) 372117-99-2P 372118-10-0P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (prepn. of heterocyclic compds. as phosphodiesterase V inhibitors preventive or therapeutic agents for various diseases due to dysfunction of signal transduction through cGMP) THERE ARE 16 CITED REFERENCES AVAILABLE REFERENCE COUNT: 16 FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L16 ANSWER 6 OF 43 CAPLUS COPYRIGHT 2002 ACS 2001:780679 CAPLUS ACCESSION NUMBER: 135:327362 DOCUMENT NUMBER: 'Nonsteroidal antiinflammatory drug (NSAID) and TITLE: NSAID derivative amyloid A.beta.42 polypeptide-lowering agents for the treatment of Alzheimer's disease , and screening methods Koo, Edward Hao Mang; Golde, Todd Eliot; INVENTOR(S): Galasko, Douglas Roger Mayo Foundation for Medical Education and PATENT ASSIGNEE(S): Research, USA SOURCE: PCT Int. Appl., 73 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE

IT

TΤ

WO 2001078721

Searcher : Shears 308-4994

WO 2001-US11956

20010412

20011025

A 1

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W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,
             CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
             LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ,
             UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD,
                                        US 2000-196617P P 20000413
PRIORITY APPLN. INFO.:
     A method is provided for preventing, delaying, or reversing the
     progression of Alzheimer's disease by administering an
     A.beta.42-lowering agent to a mammal under conditions in which
     levels of A.beta.42 are selectively reduced, levels of A.beta.38 are
     increased, and levels of A.beta.40 are unchanged. The invention
     provides methods and materials for developing and identifying
     A.beta.42-lowering agents. In addn., the invention provides methods
     for identifying agents that increase the risk of developing, or
     hasten progression of, Alzheimer's disease. The invention
     also provides compns. of A.beta.42-lowering agents and antioxidants,
     A.beta.42 lowering agents and non-selective secretase inhibitors,
     and A.beta.42 lowering agents and acetylcholinesterase inhibitors.
     The invention further provides kits contg. A.beta.42-lowering
     agents, antioxidants, non-selective secretase inhibitors, and/or
     acetylcholinesterase inhibitors as well as instructions related to
     dose regimens for A.beta.42-lowering agents, antioxidants,
     non-selective secretase inhibitors, and acetylcholinesterase
     inhibitors. The agents of the invention include nonsteroidal
     antiinflammatory drugs (NSAIDs) and NSAID derivs.
     261766-35-2 261766-36-3 261766-37-4
     261766-38-5 261766-41-0 261766-42-1
     261766-43-2
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (NSAID and NSAID deriv. amyloid A.beta.42 polypeptide-lowering
        agents for treatment of Alzheimer's disease,
        and screening methods)
                               THERE ARE 6 CITED REFERENCES AVAILABLE FOR
REFERENCE COUNT:
                         6
                               THIS RECORD. ALL CITATIONS AVAILABLE IN
                               THE RE FORMAT
L16 ANSWER 7 OF 43 CAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                         2001:661391 CAPLUS
DOCUMENT NUMBER:
                         135:210946
                         Preparation of pyridylamides as Factor Xa
                         inhibitors.
                         Zhu, Bing-yan; Zhang, Penglie; Wang, Lingyan;
INVENTOR(S):
                         Huang, Wenrong; Goldman, Erick; Li, Wenhao;
                         Zuckett, Jingmei; Song, Yonghong; Scarborough,
                         Robert
                         Cor Therapeutics, Inc., USA
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 306 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
```

TT

TITLE:

SOURCE:

Searcher : 308-4994 Shears

FAMILY ACC. NUM. COUNT: 5
PATENT INFORMATION:

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KIND
                                           APPLICATION NO.
                                                             DATE
     PATENT NO.
                            DATE
     WO 2001064642
                       A2
                            20010907
                                           WO 2001-US6247
                                                             20010228
     WO 2001064642
                      A3
                            20020502
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,
             CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
             LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ,
             UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD,
             ΤG
                                        US 2000-185746P P
                                                            20000229
PRIORITY APPLN. INFO .:
                                        US 2000-663420
                                                         A
                                                            20000915
                         MARPAT 135:210946
OTHER SOURCE(S):
    AQDEGJX [A = alkyl, cycloalkyl, NR1R2, NR1R1C(:NR3), (substituted)
     Ph, naphthyl, mono- or bicyclic heterocyclyl, etc.; R1-R3 = H,
     alkyl, alkenyl, alkynyl, cycloalkyl, (alkyl)aryl, (alkyl)heteroaryl,
     etc.; R1R2 or R2R3 = atoms to form a 3-8 membered (substituted)
     (heterocyclic) ring; Q = bond, CH2, CO, O, NR7, etc.; R7 = H, alkyl,
     (alkyl)aryl, (alkyl)heteroaryl, etc.; D = bond, (substituted) Ph,
     naphthyl, mono- or bicyclic heterocyclyl; E = bond, alkyl, S, SO,
     SO2, alkoxy, etc.; G = (substituted) alkenyl, cycloalkenyl,
     phenylene, heterocyclyl, fused cyclic system; J = bond, NR9CO, O, S,
     SO, SO2, SO2NR9, CH2, NR9, etc.; R9 = H, alkyl, (alkyl)aryl, etc.; X
     = (substituted) Ph, naphthyl, heteroaryl, fused bicyclyl), were
     prepd. as antithrombotics (no data). Thus, N-(5-bromo-2-pyridinyl)
     2-aminophenylcarboxamide (prepn. given), 4-[(2-tert-
     butylaminosulfonyl)phenyl]benzoyl chloride, and pyridine were
     stirred overnight in CH2Cl2 to give 85% N-(5-bromo-2-pyridinyl)-[2-4-
     [(2-aminosulfonyl)phenyl]phenylcarbonylamino]phenylcarboxamide.
     358659-61-7P 358659-62-8P 358659-63-9P
ΙT
     358659-64-0P 358659-65-1P 358659-66-2P
     358659-67-3P 358659-68-4P 358659-69-5P
     358659-70-8P 358659-71-9P 358659-72-0P
     358659-73-1P 358659-74-2P 358659-75-3P
     358659-76-4P 358659-77-5P 358659-78-6P
     358659-79-7P 358659-80-0P 358659-81-1P
     358659-82-2P 358659-83-3P 358659-84-4P
     358659-85-5P 358659-86-6P 358659-87-7P
     358659-88-8P
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (prepn. of pyridylamides as Factor Xa inhibitors)
L16 ANSWER 8 OF 43 CAPLUS COPYRIGHT 2002 ACS
                         2001:565010 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         135:137407
                         Preparation of 2-aminonicotinamides as
TITLE:
                         VEGF-receptor tyrosine kinase inhibitors
```

308-4994

Shears

Searcher :

INVENTOR(S): Manley, Paul William; Bold, Guido

PATENT ASSIGNEE(S): Novartis A.-G., Switz.; Novartis-Erfindungen

Verwaltungsgesellschaft m.b.H.

SOURCE: PCT Int. Appl., 66 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	ENT	NO.		KI	ND	DATE			Α	PPLI	CATI	ON NO	ο.	DATE		
WO	2001	0551	14	A	 1	2001	0802		W	0 20	01-E	P835		2001	0125	
	W:	AE,	AG,	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,
		CN,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
	•	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,
		LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,
						SD,										
		UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,
		ТJ,	TM													
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,
						FI,										
						CG,										
		TG	•	•	•	·	•	·	·	·	•	·				
RITY	APP	LN.	INFO	. :					GB 2	000-	1930		Α	2000	0127	

OTHER SOURCE(S): MARPAT 135:137407

GΙ

The title compds. [I; n = 1-6; W = O, S; R1, R3 = H, alkyl, acyl; R2 AB = (un)substituted cycloalkyl, aryl, mono- or bicyclic heteroaryl comprising one or more ring N atoms and 0-2 heteroatoms selected from O and S; R, R' = H, alkyl; X = (un)substituted aryl, mono- or bicyclic heteroaryl comprising one or more ring N atoms and 0-2heteroatoms selected from O and S] and their pharmaceutically acceptable salts, useful for therapy of a disease which responds to an inhibition of the VEGF-receptor tyrosine kinase activity (such as neoplastic disease), were prepd. and formulated. Thus, amidation of 3-aminobenzotrifluoride with 2-chloronicotinoyl chloride followed by reacting 4-pyridineethanamine with the resulting carboxamide afforded I [n = 2; R, R' = H; X = 4-pyridyl; W = 0; R1, R3 = H; R2 = 3-(F3C)C6H4]. 352227-86-2P 352227-92-0P 352228-00-3P ΙT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of 2-aminonicotinamides as VEGF-receptor tyrosine kinase

inhibitors)
REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L16 ANSWER 9 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2001:545665 CAPLUS

DOCUMENT NUMBER:

135:137515

TITLE:

Preparation of pyridines, pyrimidines,

purinones, pyrrolopyrimidinones and

pyrrolopyridinones as corticotropin releasing

factor antagonists

INVENTOR(S):

Chen, Yuhpyng Liang

PATENT ASSIGNEE(S):

Pfizer Products Inc., USA PCT Int. Appl., 130 pp.

SOURCE: PCT Int. App

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	KIND DATE				APPLICATION NO. DATE											
WO 200	WO 2001053263							W	0 20	01-I	 В4					
W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	
	CN,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KZ,	LC,	LK,	
				•		•				•		-	-	NO,		
•	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	
•	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	
		TM														
RW	: GH,															
														PT,		
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	
	TG			_								_				
US 200				1	2002	0207						-	2001			
PRIORITY AP								US 2	000-	1766	11P	Р	2000	3118		
OTHER SOURC	E(S):			MAR	PAT	135:	13/5	15								

$$R^3$$
 N
 R^6
 N
 N
 R^5
 N
 R^5
 N

The title compds. [I-III; A = CR7, N; B = NR1R2, COR2, CHR1OR2, AB etc.; G = H, O, S, etc.; Y = CH, N; Z = NH, O, S, etc.; R1 = CHO, CO(alkyl), alkyl, etc.; R2 = H, alkyl, cycloalkyl, etc.; R3 = Me, Et, F, etc.; R4 = H, alkyl, cycloalkyl, etc.; R5 = (un)substituted (hetero)aryl; R6 = H, alkyl, cycloalkyl, etc.; R16, R17 = H, OH, Me, etc.], useful in the treatment disorders including CNS and stress-related disorders, were prepd. Thus, reacting N-4-(1-ethylpropyl)-6-methyl-2-(2,4,6trimethylphenoxy)pyridine-3,4-diamine with chloroacetyl chloride in the presence of Et3N in THF afforded 91% I [A = CH; B = NHCHEt2; R3 = Me; R4 = NHCOCH2C1; Z = O; R5 = 2,4,6-Me3C6H2]. The CRF binding activities for compds. I-III, expressed as IC50 values, generally range from about 0.5 nM to 10 .mu.M.

351380-07-9P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of pyridines, pyrimidines, purinones, pyrrolopyrimidinones and pyrrolopyridinones as corticotropin

releasing factor antagonists)

REFERENCE COUNT:

THERE ARE 11 CITED REFERENCES AVAILABLE 11 FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2002 ACS L16 ANSWER 10 OF 43

ACCESSION NUMBER:

2001:453059 CAPLUS

TITLE:

135:46172

DOCUMENT NUMBER:

Preparation of N-isoxazolyl biphenylsulfonamides and related compounds as dual angiotensin II and

endothelin receptor antagonists.

INVENTOR(S):

Murugesan, Natesan; Tellew, John E.; Macor, John

E.; Gu, Zhengxiang

PATENT ASSIGNEE(S):

Bristol-Myers Squibb Co., USA

SOURCE:

PCT Int. Appl., 287 pp.

CODEN: PIXXD2

308-4994 Shears Searcher :

DOCUMENT TYPE:

Patent English

LANGUAGE:

GI

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	TENT I	.00		KI	ND	DATE			P	PPLI	CATI	ON NO	0.	DATE		
		2001								W	0 20		s337	30	2000	1213	
	0	₩:	AE, CR, HU, LT, RU, UZ, GH, CY,	AG, CU, ID, LU, SD, VN, GM, DE,	AL, CZ, IL, LV, SE, YU, KE, DK,	AM, DE, IN, MA, SG, ZA, LS, ES,	AT, DK, IS, MD, SI, ZW, MW, FI,	AU, DM, JP, MG, SK, AM, MZ, FR,	EE, KE, MK, SL, AZ, SD, GB,	ES, KG, MN, TJ, BY, SL, GR,	FI, KP, MW, TM, KG, SZ, IE,	GB, KR, MX, TR, KZ, TZ, IT,	GD, KZ, NO, TT, MD, UG, LU,	GE, LC, NZ, TZ, RU, ZW, MC,	CA, GH, LK, PL, UA, TJ, AT, NL, NE,	GM, LR, PT, UG, TM BE, PT,	HR, LS, RO, US, CH, SE,
		APP)	TG LN.		•			135:	:	US 1 US 2 US 2 US 2 US 2	999- 000-	4640 4811 5137 6043	37 97 79 22	A A A	1999 2000 2000 2000 2000	1215 0111 0225 0626	10,
OTHER		ONCE			T.T.TI	- 1.7 -		1017	-								

Ι

II

Title compds. (I; R1 = specified oxoimidazolyl, pyridoimidazolyl, pyridylamino, triazolyl, quinolinyloxy, etc.; R2 = H, halo, CHO, (halo)alkyl, cycloalkylalkyl, alkenyl, alkynyl, alkoxyalkyl, alkoxy, cyano, OH, NO2, etc.; R3 = heteroaryl; with provisos) were prepd. as dual angiotensin II and endothelin receptor antagonists for treatment of hypertension and other diseases (no data). Thus, 4-BrC6H4CH2OH was coupled with [2-[[(4,5-dimethyl-3-isoxazolyl)[(2-methoxyethoxy)methyl]amino]sulfonyl]phenyl]boronic acid to give N-(4,5-dimethyl-3-isoxazolyl)-4'-(hydroxymethyl)-N-[(2-methoxyethoxy)methyl][1,1'-biphenyl]-2-sulfonamide (66%). This was brominated to give the 4'-bromomethyl deriv. (90%), reacted with

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2-butyl-1,3-diazaspiro[4.4]non-1-en-4-one hydrochloride, and
deprotected (49% for two steps) to give II.
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254739-90-7P 254742-75-1P IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(prepn. of N-isoxazolyl biphenylsulfonamides and related compds. as dual angiotensin II and endothelin receptor antagonists)

L16 ANSWER 11 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 2001:380546 CAPLUS 134:367194

DOCUMENT NUMBER:

Preparation of novel phenylalanine derivatives

as .alpha.4-integrin inhibitors

Tanaka, Yasuhiro; Yoshimura, Toshihiko; Izawa, INVENTOR(S):

Hiroyuki; Ejima, Chieko; Kojima, Mitsuhiko; Atake, Yuko; Nakanishi, Eiji; Suzuki, Nobuyasu; Makino, Shingo; Suzuki, Manabu; Murata, Masahiro

PATENT ASSIGNEE(S): Ajinomoto Co., Inc., Japan SOURCE: PCT Int. Appl., 155 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

TITLE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA?	TENT	NO.		KI	ND	DATE			A	PPLI	CATI	ON NO	ο.	DATE		
WO	2001	0363	 76	 A	 1	20010	0525		W	0 20	 00-J	P815	 2	2000	1120	
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,
		CN,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
														ΚZ,		
		LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,
		UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,
		ТJ,	TM													
	RW:	GH,	GM,	ΚE,	LS,	MW,	ΜZ,	SD,	SL,	SZ,	ΤZ,	UG,	ZW,	ΑT,	BE,	CH,
	•	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,
		TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,
		TG														
RTTY	/ APP	T.N	TNFO	. :					JP 1	999-	3284	68	Α	1999	1118	

PRIORITY APPLN. INFO.:

JP 2000-197139 A 20000629

OTHER SOURCE(S):

MARPAT 134:367194

$$K-Z = \begin{bmatrix} G & F & E & COB \\ \hline & C & C & CH-CH_2 \end{bmatrix}$$

AB Phenylalanine derivs. represented by general formula (I) or

> 308-4994 Searcher : Shears

pharmaceutically acceptable salts thereof [wherein X represents an interat. bond, O, OSO2, N-(un) substituted NH, NHCO, NHSO2, NHCONH, or NH(CS)NH, CO; Y and Z represent each CO, SO, or SO2; A represents a specific substituted Ph group or nitrogen-contg. heterocycle such as arom.-fused pyrimidinedione or pyrimidinone, 2,4- or 2,5-imidazolidinedione, or 5-imidazolone; C represents hydrogen, lower alkyl, lower alkenyl, lower alkynyl, cyclic alkyl-lower alkyl optionally contq. heteroatoms in the ring, aryl-lower alkyl, heteroaryl-lower alkyl; D and E represent each lower alkyl, lower alkenyl, lower alkynyl, cyclic alkyl-lower alkyl optionally contg. heteroatoms in the ring, aryl-lower alkyl, heteroaryl-lower alkyl, etc. or D and E may be bonded to each other to form a ring optionally contg. 1 or 2 O, N, or S in the ring; F and G represent each hydrogen, lower alkyl, lower alkenyl, lower alkynyl, cyclic alkyl-lower alkyl optionally contg. heteroatoms in the ring, aryl-lower alkyl, heteroaryl-lower alkyl, etc. or F and G may be bonded to each other to form a ring; n is from 0 to 2; K represents OR7, NR7R8, NHNR7R8, SR7, or R7; R7 and R8 represents H, lower alkyl, etc.; and J and J' represent each hydrogen, halogeno, lower alkyl, lower alkoxy, or NO2] are prepd. These derivs. and analogs thereof show an .alpha.4 integrin inhibitory activity and are usable as remedies for various diseases relating to .alpha.4 integrin, such as inflammatory diseases related to .alpha.4 integrin-dependent adhesion process, arthritis, inflammatory intestinal diseases, systemic lupus erythematosus, multiple sclerosis, Sjoegren syndrome, psoriasis, allergy, diabetes, cardiovascular diseases, arteriosclerosis, restenosis, tumor proliferation, tumor metastasis, or transplant rejection. Thus, O-(2,6-dichlorobenzyl)-L-tyrosine bound to Wang resin was allowed to react with diethylmalonic acid, HOAt, 2-dimethylaminoisopropyl chloride hydrochloride (DIC), and N-methyl-2-pyrrolidinone (NMP) at room temp. for 16 h, washed with DMF five times, and condensed with pyrroline using HOAt, DIC, and NMP, followed by oxidn. with OsO4 in dioxane at room temp. for 16 and resin-cleavage in aq. CF3CO2H to give N-[2-[(cis-2,4dihydroxypyrrolidin-1-yl)carbonyl]-2-ethylbutanoyl]-0-(2,6dichlorobenzyl)-L-tyrosine (II). II and N-[2-[(pyrrolidin-1yl)carbonyl]-2-ethylbutanoyl]-4-(2,6-dichlorobenzoylamino)-Lphenylalanine inhibited the binding of human recombinant VCAM-1 to human B lymphoma cell line expressing integrin.alpha.4.beta.7 with IC50 of .ltoreq.0.02 .mu.mol/L.

IT 340719-28-0P 340719-29-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of novel phenylalanine derivs. as .alpha.4-integrin inhibitors)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 12 OF 43 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:319860 CAPLUS

DOCUMENT NUMBER:

134:340354

TITLE:

Preparation of anthranilamides as inhibitors of

cGMP phosphodiesterase.

INVENTOR(S):

Oku, Teruo; Sawada, Kozo; Kuroda, Akio;

Kayakiri, Natsuko; Urano, Yasuharu; Sawada,

Yuki; Mizutani, Tsuyoshi

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co., Ltd., Japan; Oku,

Noriko; Oku, Chikako; Oku, Tomohito

SOURCE: PCT Int. Appl., 105 pp.

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE: Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE _____ _____ 20001019 20010503 WO 2000-JP7308 WO 2001030745 A1 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, ΜT

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO.: AU 1999-3652 A 19991025

OTHER SOURCE (S): MARPAT 13

Ι

OTHER SOURCE(S):

MARPAT 134:340354

NHAR2

NHR3

Title compds. I; [R1 = NO2, amino, cyano, haloalkyl, acyl, halo, etc.; R2 = H, OH, alkoxy, alkyl, cycloalkyl, (substituted) aryl, heterocyclyl; A = alkylene; R3 = (substituted) heterocyclyl, CR4R5R6; R4, R5 (substituted) carbamoyl, alkyl; R4R5C = (substituted) carbocyclyl; R6 = H, alkyl], were prepd. Thus, reaction of 2-(cyclopentylamino)-5-nitrobenzoic acid with BuNH2 in DMF in the presence of 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide and 1-hydroxybenzotriazole gave N-butyl-2-(cyclopentylamino)-5-nitrobenzamide. The latter inhibited human platelet cGMP phosphodiesterase with IC50 <10 nM.

IT 337360-72-2P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of anthranilamides as inhibitors of cGMP phosphodiesterase)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 13 OF 43 CAPLUS COPYRIGHT 2002 ACS 2001:208252 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 134:252363 Preparation and effect of nitrogen-containing-TITLE: six-membered aromatic compounds as PDE V activity inhibitors Yamada, Koichiro; Matsuki, Kenji; Omori, Kenji; INVENTOR(S): Kikkawa, Kohei PATENT ASSIGNEE(S): Tanabe Seiyaku Co., Ltd., Japan PCT Int. Appl., 91 pp. SOURCE: CODEN: PIXXD2 DOCUMENT TYPE: Patent Japanese LANGUAGE: FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE ----------_____ WO 2001019802 20010322 WO 2000-JP6258 20000913 A1 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG A2 20020115 JP 2000-277652 20000913 JP 2002012587

MARPAT 134:252363

JP 1999-261852

JP 2000-130371

A 19990916

A 20000428

GI

PRIORITY APPLN. INFO.:

OTHER SOURCE(S):

AB Title compds. [I; A is an optionally substituted nitrogenous heterocyclic group; R1 is optionally substituted lower alkyl, NHQR3 (wherein R3 is an optionally substituted nitrogenous heterocyclic group; and Q is lower alkylene or a single bond), or NHR4 (wherein R4 is optionally substituted cycloalkyl); R2 is optionally substituted aryl; and either of Y and Z is CH and the other is N], pharmacol. acceptable salts are prepd. and are exhibiting an excellent selective inhibitory activity against PDE V and being useful as preventive or therapeutic drugs for erectile dysfunction (no data). Thus, the title compd. II was prepd.

ΙI

IT 330784-43-5P 330784-44-6P 330784-45-7P 330785-08-5P 330785-09-6P 330785-10-9P 330785-11-0P 330785-12-1P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL
(Biological study); PREP (Preparation); USES (Uses)
 (prepn. and effect of heteroarom. compds. as PDE V activity
inhibitors)

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FORTHIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

308-4994

L16 ANSWER 14 OF 43 CAPLUS COPYRIGHT 2002 ACS

7

ACCESSION NUMBER:

2001:152665 CAPLUS

DOCUMENT NUMBER:

134:207826

TITLE:

Preparation of substituted (aminoiminomethyl or aminomethyl)dihydrobenzofurans and benzopyrans

as factor Xa and factor IIa inhibitors

INVENTOR(S):

Burns, Christopher J.; Dankulich, William P.;

McGarry, Daniel G.; Volz, Francis A.

PATENT ASSIGNEE(S):

Aventis Pharmaceuticals Products Inc., USA

SOURCE:

PCT Int. Appl., 107 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

Searcher :

LANGUAGE:

Shears

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

	PATENT NO. K				KI	ND DATE			APPLICATION NO.						DATE			
	WO 2001014358 A WO 2001014358 A									W	20	00-1	B156	2	2000	0812		
			AE, CN, GM,	AG, CR, HR,	AL, CU, HU,	AM, CZ, ID,	AT, DE, IL, LV,	AU, DK, IN,	DM, IS,	DZ, JP,	EE, KE,	ES, KG,	FI, KP,	GB, KR,	GD, KZ,	GE, LC,	GH, LK,	
			PL,	PT, UG,	RO,	RU,	SD, VN,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	
		RW:	CY,	DE,	DK,	ES,	MW, FI, CI,	FR,	GB, GA,	GR, GN,	IE, GW,	IT, ML,	LU, MR,	MC, NE,	NL, SN,	PT, TD,	SE,	
PRIOR	ITY	APP	LN.	INFO	.:										1999 1999			
OTHER GI	SC	OURCE	(S):			MAR	PAT	134:										

The title compds. [I; n = 1 or 2; W is H or a ring system AB substituent; R is hydrogen, cyano, cycloalkyl, cycloalkenyl, heterocyclyl, fused arylcycloalkyl, fused heteroarylcycloalkyl, etc.; R1 is hydrogen, alkyl, aralkyl, heteroaralkyl, acyl, aroyl, heteroaroyl, alkoxycarbonyl, aryloxycarbonyl or heteroaryloxycarbonyl; R2 and R3 are each hydrogen, or, taken together are :NR4; R4 is hydrogen, R5O2C, HO, cyano, R5CO, HCO, lower alkyl, nitro, etc.; R5 is alkyl, aryl, heteroaryl, aralkyl, or heteroaralkyl; L1 is alkylene, alkenylene or alkynylene; L2 is absent, alkylene, alkenylene, alkynylene, alkylene-O, alkenylene-O, etc., provided that when L2 is absent, then R is not hydrogen, and Q is attached to R through a carbon atom thereof; Q is NR8', O, CO, CO2, O2C, NR8'(X1), C(X)NR8', NR8C(X1)O, etc.; provided that a nitrogen atom or oxygen atom of Q is not directly bonded to a carbon atom of L1 or L2 having a double bond or triple bond, or Q-L2-R is cycloalkyl, cycloalkenyl, heterocyclyl, fused arylcycloalkyl, fused heteroarylcycloalkyl, etc., provided that a nitrogen atom or oxygen atom of Q is not directly bonded to a carbon atom of L1 having a double bond or triple bond; X1 is O or S; R8' is hydrogen, alkyl, aralkyl, heteroaralkyl, acyl, aroyl, heteroaroyl or alkoxycarbonyl; R8 is hydrogen, alkyl, aralkyl, heteroaralkyl, acyl, aroyl or heteroaroyl; and m is 0, 1 or 2], oxides thereof, pharmaceutically acceptable salts, solvates thereof, or prodrugs thereof are prepd. These compds. inhibit the formation of simultaneously directly

inhibiting both Factor Xa and Factor IIa (thrombin) and are useful for treating pathol. conditions in a patient that may be ameliorated by administration of such compds. The pathol. conditions include venous vasculature, arterial vasculature, abnormal thrombus formation, acute myocardial infarction, unstable angina, thromboembolism, acute vessel closure assocd. with thrombolytic therapy, percutaneous transluminal coronary angioplasty, transient ischemic attacks, stroke, intermittent claudication or bypass grafting of the coronary or peripheral arteries, vessel luminal narrowing, restenosis post coronary or venous angioplasty, maintenance of vascular access patency in longterm hemodialysis patients, pathol. thrombus formation occurring in the veins of the lower extremities following abdominal, knee and hip surgery, a risk of pulmonary thromboembolism, or disseminated systemic intravascular coagulopathy occurring in vascular systems during septic shock, certain viral infections or cancer (no data). Thus, To a cooled (0.degree.) soln. of 5-(pyrid-2-yl)thiophene-2carboxylic acid and 4-methylmorpholine in CH2Cl2 is added dropwise a soln. of iso-Pr chloroformate in toluene, stirred 30 min, treated with 2-[5-(N-tert-butoxycarbonyl)carbamimidoyl-2,3dihydrobenzofuran-3-yl]ethylamine in DMF, and the reaction mixt. was allowed to warm to room temp. overnight to give 5-pyridin-2ylthiophene-2-carboxylic acid [2-[5-(N-tertbutoxycarbonyl)carbamimidoyl-2,3-dihydrobenzofuran-3-yl]ethyl]amide which was stirred with H2O and CF3CO2H in CH2Cl2 for 3 h to give 5-(pyridin-2-yl)thiophene-2-carboxylic acid [2-(5-carbamimidoyl-2,3dihydrobenzofuran-3-yl)ethyl]amide.

ΙT 328124-74-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; prepn. of substituted (aminoiminomethyl or aminomethyl)dihydrobenzofurans and -benzopyrans as inhibitors of factor Xa and factor IIa)

328123-93-9P TΤ

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of substituted (aminoiminomethyl or aminomethyl)dihydrobenzofurans and -benzopyrans as inhibitors of factor Xa and factor IIa)

L16 ANSWER 15 OF 43 CAPLUS COPYRIGHT 2002 ACS

2001:114982 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 134:173028

Cyclic amine CCR3 antagonists TITLE:

Shiota, Tatsuki; Sudoh, Masaki; Yokoyama, INVENTOR(S):

Tomonori; Muroga, Yumiko; Kamimura, Takashi;

Nakanishi, Akinobu

PATENT ASSIGNEE(S): Teijin Ltd., Japan

PCT Int. Appl., 263 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE

> 308-4994 Searcher : Shears

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WO 2000-JP5260 20000804
     WO 2001010439
                     A1
                            20010215
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,
             CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
             LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
             BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                           20020502
                                          EP 2000-950006 20000804
                       A1
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, MC, IE,
             SI, LT, LV, FI, RO, MK, CY, AL
                                         JP 1999-220864
                                                          A 19990804
PRIORITY APPLN. INFO.:
                                         WO 2000-JP5260
                                                          W 20000804
                         MARPAT 134:173028
OTHER SOURCE(S):
     Drugs contg. as the active ingredient cyclic amine derivs.
     represented by general formula (Markush's structure given),
     pharmaceutically acceptable acid addn. salts thereof or
     pharmaceutically acceptable C1-6 alkyl adducts thereof.
                                                               These drugs
     are efficacious in preventing and treating
     diseases in which CCR3 participates such as asthma and
     allergic rhinitis.
IT
     226241-69-6 308361-85-5
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (cyclic amine CCR3 antagonists as antiasthmatics and allergy
        inhibitors)
                                THERE ARE 28 CITED REFERENCES AVAILABLE
                          28
REFERENCE COUNT:
                                FOR THIS RECORD. ALL CITATIONS AVAILABLE
                                IN THE RE FORMAT
L16 ANSWER 16 OF 43 CAPLUS COPYRIGHT 2002 ACS
                          2001:63819 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                          134:131317
                          Preparation of 2-phenylaminobenzamides and
TITLE:
                          analogs as MEK inhibitors for the
                          treatment of chronic pain
                          Dixon, Alistair; Lee, Kevin; Pinnock, Robert
INVENTOR(S):
                          Denham
                         Warner-Lambert Company, USA
PATENT ASSIGNEE(S):
                          PCT Int. Appl., 132 pp.
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     WO 2001005392
                       A3
                             20010719
             AE, AG, AL, AU, BA, BB, BG, BR, BZ, CA, CN, CR, CU, CZ, DM,
             DZ, EE, GD, GE, HR, HU, ID, IL, IN, IS, JP, KP, KR, LC, LK,
             LR, LT, LV, MA, MG, MK, MN, MX, MZ, NO, NZ, PL, RO, SG, SI,
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SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ,

MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,

CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

EP 1202726 A2 20020508 EP 2000-943383 20000705

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,

PT, IE, SI, LT, LV, FI, RO, MK, CY, AL

II

PRIORITY APPLN. INFO.: US 1999-144292P P 1999071.6

WO 2000-US18347 W 20000705

OTHER SOURCE(S):

MARPAT 134:131317

GΙ

The title compds. (I) [wherein R1 = H, OH, alkyl, alkoxy, halo, CF3, AΒ or CN; R2 = H; R3, R4, and R5 = independently H, OH, halo, CF3, alkyl, alkoxy, NO2, CN, or (O or NH)m(CH2)nR9; R9 = H, OH, CO2H, or NR10R11; m = 0 or 1; n = 0-4; R10 and R11 = independently H, alkyl, or taken together with the N to which they are attached form a heterocycle; R6 = H, (cyclo)alkyl, acyl, aryl, or aralkyl; R7 = H, (cyclo)alkyl, alkenyl, alkynyl, or heterocyclyl] were prepd. using conventional and combinatorial synthetic methods for the treatment of chronic pain. For example, 2,4-difluorobenzoic acid in THF was added to a soln. of 2-amino-5-iodotoluene and Li diisopropylamide in THF/heptane/EtPh to give 4-fluoro-2-(4-iodo-2methylphenylamino)benzoic acid (47%). Treatment of the acid with O-(tetrahydro-2H-pyran-2-yl)hydroxylamine and diisopropylethylamine in THF/CH2Cl2 in the presence of PyBOP afforded the O-protected intermediate, which was dissolved in ethanolic HCl to give the title N-hydroxybenzamide (II) in 23% yield. Biol. assays indicated that MEK inhibitors exert an antiallodynic effect in CCI-induced neuropathic rats when administered intrathecally and that the antiallodynic effect correlates with the affinity of the compds. ΙT

1 212628-77-8P, 5-Chloro-N-(2-hydroxyethyl)-2-(4-iodo-2methylphenylamino)benzamide 212628-80-3P,

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4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-methylbenzamide
212628-81-4P, N-Ethyl-4-fluoro-2-(4-iodo-2-
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4-Fluoro-2-(4-iodo-2-methylphenylamino)-N, N-dimethylbenzamide
212628-83-6P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(1H-
tetrazol-5-yl)benzamide 212628-85-8P, 5-Chloro-2-(4-iodo-2-
methylphenylamino)-N, N-dimethylbenzamide 212628-86-9P,
[5-Chloro-2-(4-iodo-2-methylphenylamino)benzoylamino]acetic acid
212628-87-0P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-
propylbenzamide 212628-88-1P, 5-Bromo-N-(2-hydroxyethyl)-2-
(4-iodo-2-methylphenylamino)benzamide 212628-89-2P,
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212628-90-5P, 4-Fluoro-N-[3-[4-(2-hydroxyethyl)piperazin-1-
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dimethylbenzamide 212628-99-4P, 5-Bromo-3,4-difluoro-N-(2-
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2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-(2-
hydroxyethyl)benzamide 212629-16-8P, 4-Fluoro-2-(4-iodo-2-
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212629-17-9P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(3-
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methylphenylamino)-N-(2-thiophen-2-ylethyl)benzamide
212629-20-4P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(2-
pyrrolidin-1-ylethyl)benzamide 212629-21-5P,
2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-(2-morpholin-4-
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dimethylaminopropyl)-3,4-difluorobenzamide 212629-25-9P,
4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-pyridin-4-
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212629-27-1P, 2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-
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phenethylbenzamide 212629-31-7P, 2-(4-Bromo-2-
methylphenylamino) -3, 4-difluoro-N-(2-thiophen-2-ylethyl)benzamide
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pyridin-4-ylmethylbenzamide 212629-33-9P,
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212629-34-0P, 2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-
(2-piperidin-1-ylethyl)benzamide 212629-35-1P,
5-Chloro-N-[3-[4-(2-hydroxyethyl)piperazin-1-yl]propyl]-2-(4-iodo-2-
methylphenylamino)benzamide 212629-36-2P,
5-Fluoro-N-[3-[4-(2-hydroxyethyl)piperazin-1-yl]propyl]-2-(4-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-io
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2-(4-Iodo-2-methylphenylamino)-5-nitro-N-pyridin-4-ylmethylbenzamide
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methylphenylamino) benzamide 212629-40-8P,
5-Chloro-2-(4-iodo-2-methylphenylamino)-N-(2-piperidin-1-
ylethyl)benzamide 212629-41-9P, 5-Chloro-2-(4-iodo-2-
methylphenylamino) -N-(2-pyrrolidin-1-ylethyl)benzamide
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methylphenylamino) benzamide 212629-44-2P,
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methylphenylamino) benzamide 212629-62-4P,
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hydroxypropyl) -2-(4-iodo-2-methyl-phenylamino) benzamide
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212629-86-2P, 2-(4-Iodo-2-methylphenylamino)-5-nitro-N-(3-
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5-Bromo-N-(2-diisopropylaminoethyl)-2-(4-iodo-2-
methylphenylamino)benzamide 212629-88-4P,
 \hbox{5-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2
ylethyl)benzamide 212629-89-5P, 5-Fluoro-2-(4-iodo-2-
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212629-90-8P, N-(3-Diethylamino-2-hydroxypropyl)-5-fluoro-2-
(4-iodo-2-methylphenylamino)benzamide 212629-91-9P,
2-(4-Iodo-2-methylphenylamino)-5-nitro-N-(2-pyrrolidin-1-
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2-(4-iodo-2-methylphenylamino)-5-nitrobenzamide 212629-93-1P
, N-(2-Diisopropylaminoethyl)-5-fluoro-2-(4-iodo-2-
methylphenylamino) benzamide 212630-00-7P,
N-Cyclopropyl-5-fluoro-2-(4-iodo-2-methylphenylamino)benzamide
212630-03-0P, 5-Fluoro-N-(2-hydroxyethyl)-2-(4-iodo-2-
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sulfamoylbenzyl)benzamide 212630-07-4P,
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212630-08-5P, N-(2-Hydroxyethyl)-2-(4-iodo-2-
methylphenylamino)-5-nitrobenzamide 212630-09-6P,
2-(4-Iodo-2-methylphenylamino)-N-methyl-5-nitro-N-phenylbenzamide
212630-10-9P, 5-Chloro-N-cyclopropyl-2-(4-iodo-2-
methylphenylamino)benzamide 212630-11-0P,
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sulfamoylbenzyl)benzamide 212630-15-4P,
N-Ally1-5-chloro-2-(4-iodo-2-methylphenylamino)benzamide
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     212630-19-8P, 5-Iodo-2-(4-iodo-2-methylphenylamino)-N-(4-
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     methylphenylamino)benzamide 212630-23-4P,
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     5-Bromo-2-(4-iodo-2-methylphenylamino)-N-methyl-N-phenylbenzamide
     212630-27-8P, N-Cyclohexyl-5-iodo-2-(4-iodo-2-
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     212630-34-7P, 5-Bromo-2-(4-iodo-2-methylphenylamino)-N-(3-
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     212630-37-0P, N-Cyclohexyl-2-(4-iodo-2-methylphenylamino)-5-
     nitrobenzamide 277335-40-7P, 5-Bromo-2-(4-iodo-2-
     ethylphenylamino)-N-(2-pyrrolidin-1-ylethyl)benzamide
     321438-66-8P, N-(2-Hydroxyethyl)-2-(4-iodo-2-
     ethylphenylamino)-5-nitrobenzamide
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (prepn. of 2-phenylaminobenzamide and 2-phenylaminobenzoic acid
        MEK inhibitors by conventional and combinatorial synthetic
        methods for treatment of chronic pain)
L16 ANSWER 17 OF 43 CAPLUS COPYRIGHT 2002 ACS
                         2000:864913 CAPLUS
ACCESSION NUMBER:
                         134:4946
DOCUMENT NUMBER:
                         Thienopyrimidines, their production and use as
TITLE:
                         gonadotropin releasing hormone antagonists
                         Furuya, Shuichi; Suzuki, Nobuhiro; Choh, Nobuo;
INVENTOR(S):
                         Nara, Yoshi
                         Takeda Chemical Industries, Ltd., Japan
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 89 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                           APPLICATION NO.
                                                             DATE
     PATENT NO.
                      KIND DATE
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20000323
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     WO 2000056739
                       A1
                                           WO 2000-JP1777
         W: AE, AG, AL, AM, AU, AZ, BA, BB, BG, BR, BY, CA, CN, CR, CU,
             CZ, DM, DZ, EE, GD, GE, HR, HU, ID, IL, IN, IS, JP, KG, KR,
             KZ, LC, LK, LR, LT, LV, MA, MD, MG, MK, MN, MX, NO, NZ, PL,
             RO, RU, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, US, UZ, VN,
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         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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                       A1
                            20011219
                                            EP 2000-911308
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
             PT, IE, SI, LT, LV, FI, RO
                                                             20000426
                                            US 2000-530495
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                       В1
                            20011002
                                           US 2000-571215
                                                             20000516
     US 6340686
                       В1
                            20020122
                                                             20010921
     NO 2001004603
                       Α
                            20011126
                                            NO 2001-4603
                                                          A 19990324
PRIORITY APPLN. INFO .:
                                         JP 1999-79371
                                                          A 20000125
                                         JP 2000-18019
                                                          A3 20000323
                                         JP 2000-87051
                                         WO 2000-JP1777
                                                          W 20000323
                                         US 2000-530495
                                                          A1 20000426
OTHER SOURCE(S):
                         MARPAT 134:4946
GI
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- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- Methods for prepn. of thienopyrimidines I (R1, R2 = H, OH, AB (un) substituted C1-4 alkoxy, C1-4 alkoxy-carbonyl or C1-4 alkyl; R3 = H, halo, OH or (un) substituted C1-4 alkoxy, n = 0-5, if n = 2 then two adjacent R3 may form C1-4 alkylenedioxy; R4 = H or C1-4 alkyl; R6 = (un)substituted C1-4 alkyl or a group of the formula Q wherein R5 is hydrogen or R4 and R5 may form heterocycle); or a pharmaceutically acceptable salt thereof, having excellent GnRH-antagonizing activity, were disclosed, as well as pharmaceutical compns. for treating sex hormone-dependent diseases. Thus, compd. II [R7 = MeONHCONH (III)] was prepd. by reacting the starting amine II (R7 = NH2) with N, N'-carbonyldiimidazole followed by O-methylhydroxylamine hydrochloride. The hydrochloride salt of III demonstrated an IC50 value of 0.0001 .mu.M against binding of 125I-leuprorelin at human GnRH receptors expressed in CHO cells.
- IT 308832-00-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. of thienopyrimidines as gonadotropin releasing hormone antagonist)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 18 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2000:824101 CAPLUS

DOCUMENT NUMBER:

134:5154

TITLE:

Preparation of cyclic amine derivatives as

remedies or preventives for diseases

in association with chemokines or chemokine

receptors

INVENTOR(S):

Shiota, Tatsuki; Miyagi, Fuminori; Kamimura,

Takashi; Ohta, Tomohiro; Takano, Yasuhiro;

Horiuchi, Hideki

PATENT ASSIGNEE(S):

Teijin Limited, Japan PCT Int. Appl., 405 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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APPLICATION NO.
                                                          DATE
                     KIND DATE
    PATENT NO.
                                          _____
                                         WO 2000-JP3203 20000518
                           20001123
    WO 2000069432
                      A1
           AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN,
            CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM,
            HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL,
            PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA,
            UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ,
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH,
            CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
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                      A1 20020213
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                           20011116
                                          NO 2001-5599
                                                           20011116
    NO 2001005599
                     Α
                                       JP 1999-175856
                                                       A 19990518 '
PRIORITY APPLN. INFO.:
                                                       A 19990906
                                       JP 1999-251464
                                       WO 2000-JP3203
                                                       W 20000518
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OTHER SOURCE(S):

MARPAT 134:5154

$$\stackrel{R^1}{\triangleright} (CH_2)_{p1} - N \stackrel{(CH_2)_{m1}}{\triangleright} (CH_2)_{n} \stackrel{NCO(CH_2)_{p}}{\triangleright} \stackrel{R^4}{\mid} (CH_2)_{q} - GR6$$

$$\stackrel{R^2}{\mid} (CH_2)_{m} \stackrel{R^3}{\mid} (CH_2)_{m} \stackrel{R^3$$

AB Remedies or preventives for **diseases** in assocn. with chemokines such as MIP-1.alpha. and/or MCP-1 or chemokine receptors such as CCR1 or CCR2 contain as the active ingredient N-acyl-amino acid N-cyclic amino or N-cyclic aminoalkyl-amide derivs. represented by general formula [I; (un)substituted Ph, C3-8 cycloalkyl, arom. heterocyclyl contg. 1-3 heteroatoms selected from O, S, and/or N; R2 = H, (un)substituted C1-6 alkyl, C2-7 alkoxycarbonyl, HO, (un)substituted Ph; pl, m1 = 0-2; m = 2-4; n = 0,1; R3 = H, (un)substituted C1-6 alkyl; R4, R5 = H, OH, (un)substituted Ph or

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10/046526
     C1-6 alkyl; or R4 and R5 are combined together to form a 3- to
     5-membered hydrocarbyl; p, q = 0.1; G = CO, SO2, CO2, NR7CO, CONR7,
     NR7SO2, or SO2NR7, NHCONH, NHCSNH, NH CO2, O2CNH; R7 = H, C1-6
     alkyl; or R7 and R5 are combined together to form C2-5 alkylene; R6
     = (un) substituted Ph, C3-8 cycloalkyl, C3-6 cycloalkenyl, CH2Ph, or
     arom. heterocyclyl contg. 1-3 heteroatoms selected from O, S, and/or
     N, wherein Ph, CH2Ph, or arom. heterocyclyl group is optionally
     fused with (un) substituted benzene or arom. heterocyclyl contg. 1-3
     heteroatoms selected from O, S, and/or N], pharmaceutically
     acceptable acid-adducts thereof, or pharmaceutically acceptable C1-6
     alkyl-adducts thereof. The above diseases include
     destruction of bone or cartilage (e.g. arthritis, rheumatoid
     arthritis, osteoarthritis, osteoporosis, injury, and tumor),
     nephritis, kidney diseases, glomerulus or interstitial
     nephritis, nephrotic syndrome, demyelinating disease, or
     multiple sclerosis. Thus, N-3-ethoxybenzyl-D-methionine-N-[1-(4-
     chlorobenzyl)-4-piperazinylmethyl]amide in vitro inhibited the
     binding of human MIP-1.alpha. to THP-1 cells by >80% at 2 .mu.M.
     226241-50-5P 226241-52-7P 226241-63-0P
     226241-64-1P 226241-65-2P 226241-66-3P
     226241-67-4P 226241-68-5P 226241-69-6P
     226241-82-3P 226241-83-4P 226242-54-2P
     226243-23-8P 226243-25-0P 226243-27-2P
     226243-29-4P 226245-19-8P 308361-84-4P
     308361-85-5P
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
        (prepn. of cyclic amine derivs. as remedies or preventives for
        diseases in assocn. with chemokines or chemokine
        receptors)
REFERENCE COUNT:
                         26
                               THERE ARE 26 CITED REFERENCES AVAILABLE
                               FOR THIS RECORD. ALL CITATIONS AVAILABLE
                               IN THE RE FORMAT
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L16 ANSWER 19 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2000:756524 CAPLUS

DOCUMENT NUMBER:

TITLE:

133:321878

Preparation of cyclic protein tyrosine kinase

inhibitors

INVENTOR(S):

Das, Jagabandhu; Padmanabha, Ramesh; Chen, Ping;

Norris, Derek J.; Doweyko, Arthur M. P.;

Barrish, Joel C.; Wityak, John

PATENT ASSIGNEE(S): SOURCE:

Bristol-Myers Squibb Co., USA PCT Int. Appl., 300 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000062778			WO 2000-US9753	20000412
W: AE, A	, AM, AT	', AU, AZ, BA,	BB, BG, BR, BY, CA.	CH. CN. CR
Cu, C	, DE, DK	, DM, EE, ES,	FI, GB, GD, GE, GH,	GM HR HII
ID, II	, IN, IS	, JP, KE, KG,	KP, KR, KZ, LC, LK,	LR, LS, LT.

LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG 20020109 EP 2000-922102 20000412 EP 1169038 A1 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO BR 2000-9721 20000412 20020213 BR 2000009721 Α 20011012 20011210 NO 2001-4970 NO 2001004970 Α US 1999-129510P P 19990415 PRIORITY APPLN. INFO.: WO 2000-US9753 W 20000412 MARPAT 133:321878 OTHER SOURCE(S):

GΙ

IT

$$t-Bu \underset{H}{\bigcirc 0} \underset{N}{\overset{N}{\bigvee}} \underset{N}{\overset{Me}{\bigvee}} \underset{N}{\overset{H}{\bigvee}} \underset{Me}{\overset{Me}{\bigvee}} \underset{Me}{\overset{Me}{\bigvee}} \underset{Me}{\overset{II}{\bigvee}}$$

The title compds. [I; Q = (un) substituted 5-6 membered heteroaryl, AB aryl; Z = a single bond, R15C:CH, (CH2)m (m = 1-2); X1, X2 = H; X1 and X2 together = O, S; R1 = H, alkyl, alkenyl, etc.; R2, R3 = H, alkyl, alkenyl, etc.; R4, R5 = H, alkyl, alkenyl, etc.], useful in the treatment of protein tyrosine kinase-assocd. disorders such as immunol. and oncol. disorders (no data), were prepd. E.g., a multi-step synthesis of thiazole II was given. Compds. I are effective at 0.1-100 mg/kg/day. 302958-78-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(prepn. of cyclic protein tyrosine kinase inhibitors) THERE ARE 7 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT: THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

> 308-4994 Searcher : Shears

L16 ANSWER 20 OF 43 CAPLUS COPYRIGHT 2002 ACS

2000:721433 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 134:25114

TITLE: Aryl ureas represent a new class of

anti-trypanosomal agents

Du, Xiaohui; Hansell, Elizabeth; Engel, Juan C.; AUTHOR(S):

Caffrey, Conor R.; Cohen, Fred E.; McKerrow,

James H.

CORPORATE SÓURCE: Department of Cellular and Molecular

Pharmacology and Medicine, University of

California, San Francisco, CA, 94143-0450, USA

Chemistry & Biology (2000), 7(9), 733-742 CODEN: CBOLE2; ISSN: 1074-5521 SOURCE:

Elsevier Science Ltd. PUBLISHER:

DOCUMENT TYPE: Journal English LANGUAGE:

Background: The trypanosomal diseases including Changas' AB disease, African sleeping sickness and Nagana have a substantial impact on human and animal health worldwide. Classes of effective therapeutics are needed owing to the emergence of drug resistance as well as the toxicity of existing agents. cysteine proteases of two trypanosomes, Trypanosoma cruzi (cruzain) and Trypanosoma brucei (rhodesain), have been targeted for a structure-based drug design program as mechanistic inhibitors that target these enzymes are effective in cell-based and animal models of trypanosomal infection. Results: We have used computational methods to identify new lead scaffolds for non-covalent inhibitors of cruzain and rhodesain, have demonstrated the efficacy of these compds. in cell-based and animal assays, and have synthesized analogs to explore structure activity relationships. Nine compds. with varied scaffolds identified by DOCK4.0.1 were found to be active at concns. below 10 .mu.M against cruzain and rhodesain in enzymic studies. All hits were calcd. to have substantial hydrophobic interactions with cruzain. Two of the scaffolds, the urea scaffold and the aroyl thiourea scaffold, exhibited activity against T. cruzi in vivo and both enzymes in vitro. They also have predicted pharmacokinetic properties that meet Lipinski's "rule of These scaffolds are synthetically tractable and lend themselves to combinatorial chem. efforts. One of the compds., 5'(1-methyl-3-trifluoromethylpyrazol-5-yl)-thiophene 3'-trifluoromethylphenyl urea (D16) showed a 3.1 .mu.M IC50 against cruzain and a 3 .mu.M IC50 against rhodesain. Infected cells treated with D16 survived 22 days in culture compared with 6 days for their untreated counterparts. The mechanism of the inhibitors of these two scaffolds is confirmed to be competitive and reversible. Conclusions: The urea scaffold and the thiourea scaffold are promising leads for the development of new effective chemotherapy for trypanosomal diseases. Libraries of compds. of both scaffolds need to be synthesized and screened against a series of homologous parasitic cysteine proteases to optimize the potency of the initial leads.

TΨ 202827-87-0 312324-33-7

RL: PRP (Properties)

(aryl ureas, a new class of anti-trypanosomal agents)

REFERENCE COUNT: THERE ARE 44 CITED REFERENCES AVAILABLE 44

FOR THIS RECORD. ALL CITATIONS AVAILABLE

IN THE RE FORMAT

308-4994 Searcher : Shears

L16 ANSWER 21 OF 43 CAPLUS COPYRIGHT 2002 ACS 2000:475533 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER:

133:89332

TITLE:

Preparation of 2-(4-bromo or 4-iodo

phenylamino)benzoic acid derivatives as MEK

inhibitors for the treatment of asthma

INVENTOR(S):

Bridges, Alexander James; Dudley, David Thomas;

WO 1999-US30419 W 19991221

Mobley, James Leslie; Saltiel, Alan Robert

PATENT ASSIGNEE(S):

Warner-Lambert Company, USA

SOURCE:

PCT Int. Appl., 106 pp.

DOCUMENT TYPE:

CODEN: PIXXD2

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE PATENT NO. _____ _____ WO 1999-US30419 19991221 A2 20000713 WO 2000040235 20001109 WO 2000040235 A3 AE, AL, AU, BA, BB, BG, BR, CA, CN, CR, CU, CZ, DM, EE, GD, GE, HR, HU, ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MA, MG, MK, MN, MX, NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG A2 20011010 EP 1999-968153 19991221 EP 1140062 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO BR 1999-16785 20011023 19991221 BR 9916785 Α US 1999-115086P P 19990107 PRIORITY APPLN. INFO.:

OTHER SOURCE(S):

MARPAT 133:89332

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Br or I
$$R^{1}$$
 R^{2} R^{5} R^{4} R^{4} R^{4} R^{4} R^{5} R^{4} R^{5} R^{6} R^{7} R^{1}

The title compds. (I) [wherein R1 = H, OH, alkyl, alkoxy, halo, CF3, AB or CN; R3-R5 = independently H, OH, halo, CF3, alkyl, alkoxy, NO2, CN, or (O or NH)m-(CH2)n-R9, where R9 = H, OH, CO2H, or NR10R11; m =0 or 1; n = 0-4; R10 and R11 = H, alkyl, or taken together with the N to which they are attached form a 3-10 membered ring; Z = CO2R7, tetrazolyl, CONR6R7, CONHNR10R11, or CH2OR7; R6 and R7 = independently H, (cyclo)alkyl, alkenyl, alkynyl, acyl, (hetero)aryl, or taken together with the N to which they are attached form's 3-10 membered ring, etc.] were prepd. by std. or combinatorial synthetic methods involving the addn. of halobenzoic acids to haloanilines and

optional redn. or amidation of the acid. For example, treatment of 2-amino-5-iodotoluene in THF with LDA in THF/heptane/ethenylbenzene soln., followed by addn. of 2,4-difluorobenzoic acid in THF afforded II. In an in vitro assay, 2-(2-methyl-4-iodophenylamino)-N-hydroxy-3,4-difluoro-5bromobenzamide (PD 171984) prevented antigen-induced prodn. of interleukin 5 (IL-5) by OVA-primed splenocytes with IC50 of 117 nM. In an adoptive-transfer assay using OVA-sensitized splenocytes cultured in the presence of PD 171984, the latter inhibited BAL eosinophilic lung inflammation by 99.82% at a dose of 10 .mu.M in mice. PD 171984 also inhibited active OVA-induced eosinophilic lung inflammation in mice dosed orally at 100 .mu.M for 4 days, suppressing BAL eosinophilia by 55.26%. Thus, I are potent MEK inhibitors that are useful in the prevention and treatment of asthma. 212628-77-8P, 5-Chloro-N-(2-hydroxyethyl)-2-(4-iodo-2methylphenylamino) benzamide 212628-80-3P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-methylbenzamide 212628-81-4P, N-Ethyl-4-fluoro-2-(4-iodo-2methylphenylamino) benzamide 212628-82-5P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N, N-dimethylbenzamide 212628-83-6P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(1Htetrazol-5-yl)-benzamide 212628-85-8P, 5-Chloro-2-(4-iodo-2-methylphenylamino)-N,N-dimethylbenzamide 212628-86-9P, [[5-Chloro-2-(4-iodo-2methylphenylamino)benzoyl]amino]acetic acid 212628-87-0P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-propylbenzamide 212628-88-1P, 5-Bromo-N-(2-hydroxyethyl)-2-(4-iodo-2methylphenylamino) benzamide 212628-89-2P, N, N-Diethyl-4-fluoro-2-(4-iodo-2-methylphenylamino)benzamide 212628-90-5P, 4-Fluoro-N-[3-[4-(2-hydroxyethyl)piperazin-1yl]propyl]-2-(4-iodo-2-methylphenylamino)benzamide 212628-91-6P, N, N-Diethyl-2-(4-iodo-2-methylphenylamino)-5nitrobenzamide 212628-92-7P, N-Butyl-4-fluoro-2-(4-iodo-2methylphenylamino) benzamide 212628-93-8P, 5-Chloro-N, N-diethyl-2-(4-iodo-2-methylphenylamino)benzamide 212628-94-9P, 5-Bromo-2-(4-iodo-2-methylphenylamino)-N, Ndimethylbenzamide 212628-99-4P, 5-Bromo-3,4-difluoro-N-(2hydroxyethyl) -2-(4-iodo-2-methylphenylamino) benzamide 212629-00-0P, N-(2,3-Dihydroxypropyl)-3,4-difluoro-2-(4-iodo-2-methylphenylamino)benzamide 212629-01-1P, 5-Bromo-3, 4-difluoro-2-(4-iodo-2-methylphenylamino)-N-(2-piperidin-1ylethyl)benzamide 212629-02-2P, 3,4-Difluoro-N-(2hydroxyethyl)-2-(4-iodo-2-methylphenylamino)benzamide 212629-03-3P, N-(2,3-Dihydroxypropyl)-4-fluoro-2-(4-iodo-2methylphenylamino)benzamide 212629-04-4P, 3,4-Difluoro-N-(3-hydroxypropyl)-2-(4-iodo-2methylphenylamino)benzamide 212629-05-5P, 5-Bromo-3,4-difluoro-2-(4-iodo-2-methylphenylamino)-N-(2-pyrrolidin-1-ylethyl)benzamide 212629-06-6P, 5-Bromo-3,4-difluoro-2-(4-iodo-2-methylphenylamino)-N-(2-pyridin-4-ylethyl)benzamide 212629-07-7P, 4-Fluoro-N-(2-hydroxyethyl)-2-(4-iodo-2methylphenylamino) benzamide 212629-08-8P, 5-Bromo-N-(3-dimethylaminopropyl)-3,4-difluoro-2-(4-iodo-2methylphenylamino)benzamide 212629-09-9P, 5-Bromo-3,4-difluoro-2-(4-iodo-2-methylphenylamino)-N-(2-morpholin-4ylethyl)benzamide 212629-10-2P, 3,4-Difluoro-2-(4-iodo-2methylphenylamino) -N-(2-morpholin-4-ylethyl)benzamide

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212629-11-3P, 3,4-Difluoro-2-(4-iodo-2-methylphenylamino)-N-
(2-pyrrolidin-1-ylethyl)benzamide 212629-12-4P,
3,4-Difluoro-2-(4-iodo-2-methylphenylamino)-N-(2-pyridin-4-
ylethyl)benzamide 212629-13-5P, N-(3-Dimethylaminopropyl)-
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212629-14-6P, N-Benzyl-4-fluoro-2-(4-iodo-2-
methylphenylamino)benzamide 212629-15-7P,
2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-(2-
hydroxyethyl)benzamide 212629-16-8P, 4-Fluoro-2-(4-iodo-2-
methylphenylamino)-N-(2-morpholin-4-ylethyl)benzamide
212629-17-9P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(3-
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3,4-Difluoro-2-(4-iodo-2-methylphenylamino)-N-(3-piperidin-1-
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212629-20-4P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(2-
pyrrolidin-1-ylethyl)benzamide 212629-21-5P,
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212629=23-7P, 3,4-Difluoro-2-(4-iodo-2-methylphenylamino)-N-
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difluorobenzamide 212629-25-9P, 4-Fluoro-2-(4-iodo-2-
methylphenylamino) -N-pyridin-4-ylmethylbenzamide
212629-26-0P, 4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(2-
pyridin-4-ylethyl)benzamide 212629-27-1P,
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212629-29-3P, 2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-
(2-pyrrolidin-1-ylethyl)benzamide 212629-30-6P,
4-Fluoro-2-(4-iodo-2-methylphenylamino)-N-phenethylbenzamide
212629-31-7P, 2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-
(2-thiophen-2-ylethyl)benzamide 212629-32-8P,
2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-pyridin-4-
vlmethylbenzamide 212629-33-9P, 2-(4-Bromo-2-
methylphenylamino)-3,4-difluoro-N-phenethylbenzamide
212629-34-0P, 2-(4-Bromo-2-methylphenylamino)-3,4-difluoro-N-
(2-piperidin-1-ylethyl)benzamide 212629-35-1P,
5-Chloro-N-[3-[4-(2-hydroxyethyl)piperazin-1-yl]-propyl]-2-(4-iodo-2-
methylphenylamino) benzamide 212629-36-2P,
5-Fluoro-N-[3-[4-(2-hydroxyethyl)piperazin-1-yl]-propyl]-2-(4-iodo-2-
methylphenylamino) benzamide 212629-37-3P,
2-(4-Iodo-2-methylphenylamino)-5-nitro-N-pyridin-4-ylmethylbenzamide
212629-38-4P, 5-Bromo-N-[3-[4-(2-hydroxyethyl)piperazin-1-
yl]-propyl]-2-(4-iodo-2-methylphenylamino)benzamide
212629-39-5P, 5-Chloro-N-(2-diethylaminoethyl)-2-(4-iodo-2-
methylphenylamino) benzamide 212629-40-8P,
5-Chloro-2-(4-iodo-2-methylphenylamino)-N-(2-piperidin-1-
ylethyl)benzamide 212629-41-9P, 5-Chloro-2-(4-iodo-2-
methylphenylamino) -N-(2-pyrrolidin-1-ylethyl)benzamide
212629-42-0P, 5-Bromo-N-(2-diethylaminoethyl)-2-(4-iodo-2-
methylphenylamino) benzamide 212629-43-1P,
N-[2-[Bis-(2-hydroxyethyl)amino]ethyl]-5-chloro-2-(4-iodo-2-
methylphenylamino) benzamide 212629-44-2P,
N-[2-[Bis-(2-hydroxyethyl)amino]ethyl]-5-bromo-2-(4-iodo-2-
methylphenylamino)benzamide 212629-46-4P,
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N-[3-[4-(2-Hydroxyethyl)piperazin-1-yl]-propyl]-2-(4-iodo-2-
methylphenylamino) benzamide 212629-47-5P,
5-Fluoro-2-(4-iodo-2-methylphenylamino)-N-pyridin-4-
ylmethylbenzamide 212629-48-6P, 5-Bromo-2-(4-iodo-2-
methylphenylamino) -N-(2-pyrrolidin-1-ylethyl)benzamide
212629-50-0P, 5-Bromo-2-(4-iodo-2-methylphenylamino)-N-(2-
piperidin-1-ylethyl)benzamide 212629-52-2P,
5-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(2-pyrrolidin-1-
ylethyl)benzamide 212629-54-4P, 5-Chloro-N-(3-
dimethylaminopropyl)-2-(4-iodo-2-methylphenylamino)benzamide
212629-56-6P, N-[2-[Bis-(2-hydroxyethyl)amino]ethyl]-5-
fluoro-2-(4-iodo-2-methylphenylamino)benzamide 212629-58-8P
   5-Chloro-N-(3-hydroxypropyl)-2-(4-iodo-2-
methylphenylamino) benzamide 212629-60-2P,
5-Chloro-N-[3-(N, N-diethylamino)-2-hydroxypropyl]-2-(4-iodo-2-
methylphenylamino) benzamide 212629-62-4P,
5-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(2-piperidin-1-
ylethyl)benzamide 212629-64-6P, 5-Bromo-N-(3-
hydroxypropyl)-2-(4-iodo-2-methylphenylamino)benzamide
212629-66-8P, 5-Bromo-2-(4-iodo-2-methylphenylamino)-N-(3-
piperidin=1-ylpropyl)benzamide 212629-68-0P,
N-[2-[Bis-(2-hydroxyethyl)amino]ethyl]-2-(4-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-2-iodo-
methylphenylamino)-5-nitrobenzamide 212629-69-1P,
\hbox{5-Chloro-2-(4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)-N-(2-morpholin-4-iodo-2-methylphenylamino)}\\
ylethyl)benzamide 212629-71-5P, 5-Chloro-N-(3-
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212629-73-7P, 5-Chloro-N-(2-diisopropylaminoethyl)-2-(4-iodo-
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212629-92-0P 212629-93-1P, N-(2-
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methylphenylamino) benzamide 212630-00-7P,
N-Cyclopropyl-5-fluoro-2-(4-iodo-2-methylphenylamino)benzamide
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212630-03-0P, 5-Fluoro-N-(2-hydroxyethyl)-2-(4-iodo-2-
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2-(4-Iodo-2-methylphenylamino)-5-nitro-N-(4-
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5-Fluoro-2-(4-iodo-2-methylphenylamino)-N-(4-
sulfamoylbenzyl)benzamide 212630-15-4P,
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iodo-2-methylphenylamino)benzamide 212630-18-7P,
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sulfamoylbenzyl)benzamide 212630-21-2P,
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212630-27-8P, N-Cyclohexyl-5-iodo-2-(4-iodo-2-
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212630-31-4P, 5-Iodo-2-(4-iodo-2-methylphenylamino)-N-methyl-
N-phenylbenzamide 212630-32-5P, N-Cyclohexyl-5-fluoro-2-(4-
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212630-34-7P, 5-Bromo-2-(4-iodo-2-methylphenylamino)-N-(3-
methylbenzyl)benzamide 212630-35-8P, 5-Bromo-N-cyclohexyl-
2-(4-iodo-2-methylphenylamino)benzamide 212630-36-9P,
5-Chloro-2-(4-iodo-2-methylphenylamino)-N-(3-methylbenzyl)benzamide
212630-37-0P, N-Cyclohexyl-2-(4-iodo-2-methylphenylamino)-5-
nitrobenzamide 277315-10-3P
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological study, unclassified); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
   (prepn. of 2-(4-bromo or 4-iodo phenylamino)benzoic acid derivs.
   as MEK inhibitors by addn. of halobenzoic acids to haloanilines
   and optional redn. or amidation of the acid)
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DOCUMENT NUMBER: 132:334364

TITLE:

Preparation of anthranilic acid amides as vascular endothelial growth factor receptor inhibitors.

INVENTOR(S):

Huth, Andreas; Seidelmann, Dieter; Thierauch, Karl-Heinz; Bold, Guido; Manley, Paul William; Furet, Pascal; Wood, Jeanette Marjorie; Mestan, Jurgen; Bruggen, Jose; Ferrari, Stefano; Kruger, Martin; Ottow, Eckhard; Menrad, Andreas; Schirner, Michael

PATENT ASSIGNEE(S):

Schering Aktiengesellschaft, Germany; Novartis

Aktiengesellschaft

SOURCE: PCT Int. Appl., 96 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent German

LANGUAGE: GEFAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PF	PATENT NO.					DATE				APPLI	CATI	ON NO	٥.	DATE		
	2000									wo 19	99-E	P847	8	1999	1109	
								BA,	ВВ	, BG,	BR,	BY,	CA,	CH,	CN,	CR,
		CU,	CZ,	DE,	DK,	DM,	EE,	ES,	FI	, GB,	GD,	GE,	GH,	GM,	HR,	HU,
		ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP	, KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,
				•	•	•		•		, MX,						
										, TR,					US,	UZ,
										, KZ,						
	RW:									, TZ,						
•										, IT,						BF,
										, ML,						
	1991									DE 19	99-1	9910.	396	19990	J303	
DE	1991	0396		C	2	2001	1213									
	9915															
. EF														1999:		
	R:		-						GB	, GR,	IT,	LI,	LU,	NL,	SE,	MC,
			ΙE,													
	2001															
PRIORIT	Y APP	LN.	INFO	.:												
										1999-				19990		
										1999-	EP84	78	W	1999	1109	
OTHER S	COURCE	(S):			MAR	PAT	132:	3343	64							

OTHER SOURCE(S):

MARPAT 132:334364

R5 AZR1

NG XYR3

AΒ

Ι

Title compds. [I; A = NR2; W = O, S, H2, NR8; Z = NR10, N, NR10(CH2)q, alkyl, etc.; q = 1-6; AZR1 = tetrahydroisoquinolinyl,

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indazolyl, 5-chloroindolyl, etc.; R1 = (substituted) aryl,
     heteroaryl; R2 = H, alkyl; R3 = (substituted) mono- or bicyclic
     aryl, heteroaryl; R4-R7 = H, halo, (substituted) alkoxy, alkyl,
     carboxyalkyl; R5R6 = dioxetanyl; R8, R10 = H, alkyl]. Thus, Me
     N-(4-pyridylmethyl)anthranilate (prepn. given) was stirred with
     Ph(CH2)3NH2 and Me3Al were stirred in PhMe to give
     N-(3-phenylprop-1-yl)-N2-(4-pyridylmethyl)anthranilamide.
     latter inhibited VEGFR I with IC50 = 0.05 .mu.M.
     267891-62-3P
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); RCT (Reactant); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); RACT (Reactant or reagent); USES (Uses)
        (prepn. of anthranilic acid amides as VEGF receptor inhibitors)
     267891-61-2P 267891-63-4P
     RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (prepn. of anthranilic acid amides as VEGF receptor inhibitors)
L16 ANSWER 23 OF 43 CAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                         2000:314687 CAPLUS
DOCUMENT NUMBER:
                         132:334454
                         Preparation of 2-amino-thiazole derivatives as
TITLE:
                         antitumor agents
                         Pevarello, Paolo; Amici, Raffaella; Traquandi,
INVENTOR(S):
                         Gabriella; Villa, Manuela; Vulpetti, Anna;
                         Isacchi, Antonella
                         Pharmacia & Upjohn S.p.A., Italy
PATENT ASSIGNEE(S):
SOURCE:
                         PCT Int. Appl., 115 pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND DATE
                                           APPLICATION NO.
                                                            DATE
                            ____
                                           ______
                                                           19991027
                                           WO 1999-EP8306
     WO 2000026202
                       A1
                            20000511
            AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, EE, GD, GE, HR, HU,
             ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MG, MK, MN,
             MX, NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN,
             YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
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ΙT

IT

GΙ

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DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF,
                     CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
             BJ, CF,
                                           EP 1999-955931
                                                            19991027
    EP 1124810
                       A1
                            20010822
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
             PT, IE, SI, LT, LV, FI, RO
                                           BR 1999-14958
                                                             19991027
    BR 9914958
                       Α
                            20011218
                                           NO 2001-2057
                                                             20010426
    NO 2001002057
                       Α
                            20010628
                                                         A 19981030
PRIORITY APPLN. INFO.:
                                        GB 1998-23871
                                                         A 19981030
                                        US 1998-823871
                                        WO 1999-EP8306
                                                         W 19991027
OTHER SOURCE(S):
                         MARPAT 132:334454
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308-4994 Searcher : Shears

$$\mathbb{R} \xrightarrow{\mathbb{N}} \mathbb{N} \xrightarrow{\mathbb{N}} \mathbb{R}^{1}$$

AB The title compds. [I; R = halo, NO2, (un) substituted amino NH2, etc.; R1 = alkyl, alkenyl, 3-6 membered carbocycle, etc.], useful for treating cell proliferative disorders assocd. with an altered cell dependent kinase activity such as cancer, Alzheimer's disease, viral infections, autoimmune diseases or neurodegenerative disorders, were prepd. E.g., thiazole I [R = iso-Pr; R1 = 4-Me2NC6H4CH2] showed Ki of 0.1 .mu.M against cdk2/cyclin A complex.

IT 267656-17-7P 267656-22-4P
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological study, unclassified); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of 2-amino-thiazole derivs. as antitumor agents)
REFERENCE COUNT: 21 THERE ARE 21 CITED REFERENCES AVAILABLE
FOR THIS RECORD. ALL CITATIONS AVAILABLE
IN THE RE FORMAT

L16 ANSWER 24 OF 43 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2000:314533 CAPLUS

DOCUMENT NUMBER:

132:334285

TITLE:

Preparation of phenyloxoazapropylcycloalkane derivatives and analogs as potassium channel

inhibitors

INVENTOR(S):

Baker, Robert K.; Chee, Jennifer; Bao, Jianming; Garcia, Maria L.; Kaczorowski, Gregory J.; Kotliar, Andrew; Kayser, Frank; Liu, Chou Juitsai; Miao, Shouwu; Rupprecht, Kathleen M.; Parsons, William H.; Schmalhofer, William A.;

Claiborne, Christopher F.; Liverton, Nigel; Claremon, David A.; Thompson, Wayne J.

PATENT ASSIGNEE(S):

SOURCE:

Merck & Co., Inc., USA PCT Int. Appl., 243 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA?	TENT	NO.		KI	ND I	DATE			A	PPLI	CATIO	ои ис	٥.	DATE		
WO	WO 2000025770 W: AE, AL			A.	1 :	2000	0511		W	19	99-U	S249	49	1999	1026	
	W:	ΑE,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,
		CU,	CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,
		ID,	IL,	IN,	IS,	JP,	KE,	KG,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,
		LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,
		SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	ΤZ,	UA,	UG,	US,	UZ,	VN,

YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,

DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF,

BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

EP 1143965 A1 20011017 EP 1999-955159 19991026

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,

PT, IE, SI, LT, LV, FI, RO

PRIORITY APPLN. INFO.: US 1998-106416P P 19981030

WO 1999-US24949 W 19991026

OTHER SOURCE(S):

MARPAT 132:334285

GΙ

$$\begin{array}{c|c}
R^1 & R^{10} \\
N - CO \\
CH_2 & R^7
\end{array}$$

$$\begin{array}{c|c}
R^6 \\
R^7 \\
R^3 & R^5
\end{array}$$

AB The title compds. I [T1 = (CH2)x; T2 = (CH2)y; dotted line indicates a single bond or double bond; x, y = 0 - 2; R1, R2, R6, R7 = halo, hydroxy, alkyl, etc.; R3, R4 = H, cyano, nitro, etc.; further details on R3 and R4 are given; R5 = H, halo, hydoxy, etc.; further details on R3 and R5 are given; R10 = H, etc.], useful as potassium channel inhibitors (no data), are prepd. I are useful in the treatment of autoimmune disorders, cardiac arrhythmias (no data), etc. Formulations are given.

Ι

IT 267405-06-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. and effect of phenyloxoazapropylcycloalkane derivs. and analogs with potassium channel inhibiting activity)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 25 OF 43 CAPLUS COPYRIGHT 2002 ACS

2

ACCESSION NUMBER: 2000:210152 CAPLUS

DOCUMENT NUMBER: 132:251068

TITLE: Preparation of N-phenylthiopheneimidamides and

analogs as NO synthase inhibitors and oxygen

scavengers

INVENTOR(S): Bigg, Dennis; Chabrier De Lassauniere,

Pierre-Etienne; Auvin, Serge; Harnett, Jeremiah;

Ulibarri, Gerard

PATENT ASSIGNEE(S): Societe De Conseils De Recherches Et

D'Applications Scientifiques (S.C.R.A.S, Fr.

SOURCE:

è

PCT Int. Appl., 74 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO.				ND	DATE			A	PPLI			ο.	DATE		
WO	2000	0171	91	A2					W				1	19990	0922	
""		ΑE,	AL,	AM,	AT,	ΑU,	ΑZ,							CH, HU,		
		IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LS,	LT,	LU, SE,	LV,	MD,
		AM,	AZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM				YU,		
	RW:	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙT,	LU,	MC,	NL,	BE, PT,	SE,	
	2784	678 ·		A	L	2000	0421		F	R 19	98-1	1867			0923	
BR	9956 9913 1115	399		Α		2001	0703		Е	R 19	99-1	3899		19990	0922	
Er		AT,	BE,	CH,	DE,		ES,	FR,						NL,		MC,
NO PRIORITY	2001 (APP	0014	78	Α		2001	0322		N FR 1	O 20 998-	01-1 1186	478 7	А	20010 19980	0322 0923	
OTHER SO						PAT			WO 1					1999		

GΙ

R1Z1Z2ZNCRNH2 [I; R = CH2NO2, alkyl, (hetero)aryl, (di)(alkyl)amino, AΒ etc.; R1 = (un) substituted anilinophenyl, -phenoxyphenyl, -C-attached carbazolyl, etc.; Z = bond or phenylene; Z1 = bond, O, S, NH, CH2NH, CO, CONH, etc.; Z2 = bond, O, NH, oxyalkylene, (heteroatom-interrupted) alkylene, etc.] were prepd. Thus, 4-(H2N)C6H4NHPh was amidated by Me 2-thiophenethiocarboximidate hydroiodide to give title compd. II.HI. Data for biol. activity of I were given.

II

262447-33-6P 262447-34-7P IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of N-phenylthiopheneimidamides and analogs as NO synthase inhibitors and oxygen scavengers)

> Shears 308-4994 Searcher :

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L16 ANSWER 26 OF 43 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2000:34745 CAPLUS DOCUMENT NUMBER: 132:93309
TITLE: Preparation of N-isoxazo
```

ITLE: Preparation of N-isoxazolyl biphenylsulfonamides and related compounds as dual angiotensin II and

endothelin receptor antagonists.

INVENTOR(S): Murugesan, Natesan; Tellew, John E.; Macor, John

E.; Gu, Zhengxiang

PATENT ASSIGNEE(S): Bristol-Myers Squibb Co., USA

SOURCE: PCT Int. Appl., 283 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	NO.	KIND	DATE							DATE		
	001300		200001	12		100				1000	7701	
												0.0
W:			U, AZ, E									
	DE, DK,	EE, E	S, FI, G	SB, GE,	GH,	GM,	HU,	ID,	IL,	IN,	IS,	JP,
	KE, KG	KP, K	R, KZ, I	C, LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,
•	MN, MW,	MX, N	O, NZ, E	PL, PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,
			T, UA, U									
	KZ, MD	RU, T	J, TM									•
RW:	GH, GM	KE, L	S, MW, S	SD, SL,	SZ,	UG,	ZW,	ΑT,	ВĒ,	CH,	CY,	DE,
			R, GB, G									
	CF, CG	CI, C	M, GA, G	SN, GW,	ML,	MR,	NE,	SN,	TD,	TG		
AU 9950	888	A1	200001	124	ΑU	J [.] 199	9-50	888		1999	0701	
EP 1094	816	A1	200105	502	EF	9 199	9-93	3540	5	1999	0701	
R:	AT, BE,	CH, D	E, DK, E	ES, FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,
	PT, IE,	SI, L	T, LV, E	ri, RO								
BR 9911	621	Α	200110	16	BF	२ 199	9-11	621		1999	0701	
LT 4854		В	200111	126	LI	r 200	0-12	23		2000	1222	
NO 2001	000062	Α	200103	305	NC	200	1-62	2		2001		
PRIORITY APP	LN. INFO).:	•		US 19	98-9	1847	7 P	Р	19980	0706	
					WO 19					1999		
OTHER SOURCE	(S):	М	ARPAT 13	32:9330	9							

GI

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Title compds. (I; R1 = specified oxoimidazolyl, pyridoimidazolyl,
AB
           pyridylamino, triazolyl, quinolinyloxy, etc.; R2 = H, halo, CHO,
           alkyl, haloalkyl, cycloalkylalkyl, alkenyl, alkynyl, alkoxyalkyl,
           alkoxy, cyano, OH, NO2, etc.; R3 = heteroaryl; with provisos), were
           prepd. as dual angiotensin II and endothelin receptor antagonists
           (no data). Thus, 4-BrC6H4CH2OH was coupled with
           [2-[[(4,5-dimethyl-3-isoxazolyl)[(2-methoxyethoxy)methyl]amino]sulfo
           nyl]phenyl]boronic acid to give N-(4,5-dimethyl-3-isoxazolyl)-4'-
           (hydroxymethyl)-N-[(2-methoxyethoxy)methyl][1,1'-biphenyl]-2-
           sulfonamide. This was brominated to give 4'-bromomethyl-N-(4,5-
           dimethyl-3-isoxazolyl)-N-[(2-methoxyethoxy)methyl][1,1'-biphenyl]-2-
           sulfonamide, which reacted with 2-butyl-1,3-diazaspiro[4.4]non-1-en-
           4-one hydrochloride followed by deprotection to give
           4'-[(2-buty)-4-oxo-1, 3-diazaspiro[4.4]non-1-en-3-y1)methy1]-N-(4,5-buty)-4-oxo-1, 3-diazaspiro[4.4]non-1-en-3-y1)methy1]-N-(4,5-buty)-4-oxo-1, 3-diazaspiro[4.4]non-1-en-3-y1)methy1]-N-(4,5-buty)-4-oxo-1, 3-diazaspiro[4.4]non-1-en-3-y1)methy1]-N-(4,5-buty)-4-oxo-1, 3-diazaspiro[4.4]non-1-en-3-y1)methy1]-N-(4,5-buty)-4-oxo-1, 3-diazaspiro[4.4]non-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1-1-en-3-y1)methy1]-N-(4,5-buty)-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y1)methy1-1-en-3-y10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-3-w10-en-
           dimethyl-3-isoxazolyl)[1,1'-biphenyl]-2-sulfonamide.
           254739-90-7P 254742-75-1P
TΤ
           RL: BAC (Biological activity or effector, except adverse); BSU
           (Biological study, unclassified); SPN (Synthetic preparation); THU
            (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
            (Uses)
                  (prepn. of N-isoxazolyl biphenylsulfonamides and related compds.
                  as dual angiotensin II and endothelin receptor antagonists)
                                                                     THERE ARE 4 CITED REFERENCES AVAILABLE FOR
REFERENCE COUNT:
                                                        4
                                                                     THIS RECORD. ALL CITATIONS AVAILABLE IN
                                                                     THE RE FORMAT
L16 ANSWER 27 OF 43 CAPLUS COPYRIGHT 2002 ACS
                                                        1999:795681 CAPLUS
ACCESSION NUMBER:
                                                        132:35606
DOCUMENT NUMBER:
                                                        Preparation of multibinding piperidinylindole
TITLE:
                                                        derivatives as therapeutic agents that
```

modulate 5-HT receptors

INVENTOR(S):

Marquess, Daniel; Griffin, John H.; Choi,

Seok-Ki

PATENT ASSIGNEE(S):

Advanced Medicine, Inc., USA

SOURCE:

PCT Int. Appl., 190 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

24

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	ENT	NO.		KI	ND I	DATE			Al	PPLIC	CATIO	ON NO).	DATE		
WO	9964	044		A.	1 :	1999:	1216		W	199	99-US	51275	51	19990	0607	
	W:	ΑE,	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,
		CZ,	DE,	DK,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	ΗU,	ID,	IL,
		IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,
		MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,
		SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW,
		AM,	ΑZ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM						
	RW:	GH,	GM,	ΚE,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,
		DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
						GA,										
CA	2318	894		ΑZ	A	1999:	1216		C	A 199	99-23	31889	94	1999	0604	
ΑU	9945	435		A:	1	1999:	1230		Αl	U 199	99-45	5435		1999	0604	
ΕP	1003	540		A:	1 .	2000	0531		E	P 199	99-92	2834	4	1999	0604	
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,

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PT, IE, FI
                                       CA 1999-2318055
                                                        19990607
                        19991216
CA 2318055
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PRIORITY APPLN. INFO.:
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OTHER SOURCE(S):
                       MARPAT 132:35606
GΙ
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$$N-CH_2-CH_2-N$$
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 F
 F
 $CO-NH$
 II

Novel multibinding piperidinylindole compds, LpXq [where L = aAΒ ligand capable of binding to a 5-HT receptor; X = a linker; p = 2-10; q = 1-2], that modulate 5-HT receptors are disclosed. Preferred ligands are of formula I [where R3 and R5 = independently point of attachment of the linker, H, alkyl, heterocyclic, heteroaryl(alkyl), amidoalkyl, (di)alkylaminosulfonylalkyl, arylsulfonylalkyl, heterocyclosulfonylalkyl, arylcarbonylamino, alkylsulfonamido, or alkylsufonylalkyl]. Over 140 multibinding compds., formed from two piperidinylindole derivs. and a difunctional linker, were prepd. For example, condensation of 5-(4-fluorobenzoyl)amino-3-(piperidin-4-yl)-1H-indole with 1,2-dibromoethane at 72.degree. in DMF, after workup and chromatog., yielded the dimer II. Compds. of this invention are useful in the treatment of migraine, headache, itch, motion sickness, depression, emesis, memory loss, anxiolytic disorders, obesity, gastrointestinal disorders, and irritable bowel syndrome (no data). The multibinding compds. provide greater biol. and/or therapeutic effects than the aggregate of the unlinked ligands due to their multibinding properties (no data). Combinatorial arrays, methods of synthesis, and methods of assaying the dimeric and multimeric compds. are also embodied by the invention.

IT 252355-17-2P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(target compd.; prepn. of multibinding piperidinylindole derivs. as **therapeutic** agents that modulate 5-HT receptors and are useful for the **treatment** of migraine)

REFERENCE COUNT:

THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 28 OF 43 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1999:409260 CAPLUS

10

DOCUMENT NUMBER:

131:73440

Preparation of aromatic amide derivatives as ACC TITLE:

inhibitor

Igawa, Hiroshi; Nishimura, Masato; Okada, Keiji; INVENTOR(S):

Nakamura, Takashi

Fujirebio, Inc., Japan PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 72 pp. SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE DATE PATENT NO. KIND _____ -----JP 1998-270721 19980925 JP 11171848 19990629 A2 JP 1997-277942 19970926 PRIORITY APPLN. INFO.:

MARPAT 131:73440 OTHER SOURCE(S):

GI

Title compds. [I; R = 3-CF3C6H4, C6H5(CH2)2, C6H5, CH3(CH2)5, AΒ CH3(CH2)3, CH3(CH2)2, CH3CH2, CH3, C6H5(CH2)3, etc.; R1 = H, CH3(CH2)4, 5-CH3(CH2)5CC, 5-CH3CH2CC, 5-(CH3)3CCC, 4-C6H5CH2O, 4-C6H5CC, 3-C6H5CC, 3-C6H5CC, 3-(4-NO2C6H4)CC, 3-(4-NCC6H4)CC, 3-(4-HOC6H4)CC, etc.; R2 = 5-OH, 5-Cl, 5-OMe, 5-Me, 5-Br, etc.; R3 = H, CH3, etc.; R4 = CO2H, AcNHSO2, CH3(CH2)4CONHSO2, 4-CF3C6H4CONHSO2, PHCONHSO2, (CH3)3CONHSO2, CH3(CH2)2NHCONHSO2, etc.; X = CH, N; dotted bond = single, double] are prepd. and tested as ACC (acetyl-CoA carboxylase) inhibitors in treatment of lipids oxidn. related diseases, such as myocardial infarction, cerebral infarction, and diabetes. The title compd. I (R = 3-CF3C6H4; R1 = H; R2 = H; R3 = H; X = CH; dotted bonds weredouble bonds) was prepd. with 72% yield from 3-EtO2CC6H4NH2 and

3-(2-HO2CC6H4NH)C6H4CF3. 228580-56-1P 228580-57-2P 228580-59-4P IT228580-72-1P 228580-98-1P 228581-26-8P 228581-28-0P 228581-31-5P 228581-32-6P 228581-34-8P 228581-35-9P 228581-36-0P 228581-38-2P 228581-39-3P 228581-40-6P 228581-42-8P 228581-43-9P 228581-44-0P 228581-57-5P 228581-58-6P 228581-59-7P

228581-60-0P 228581-61-1P 228581-62-2P

228581-63-3P 228581-64-4P 228581-65-5P

228581-66-6P 228581-68-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

> 308-4994 Shears Searcher :

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(prepn. of arom. amide derivs. as ACC inhibitor)
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     228580-62-9P 228580-89-0P 228581-54-2P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
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L16 ANSWER 29 OF 43 CAPLUS COPYRIGHT 2002 ACS
                         1999:350650 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         131:18925
                         Preparation of cyclic amine derivatives for
TITLE:
                         inhibition of the action of chemokines such as
                         MIP-1.alpha. and/or MCP-1 on target cells
                         Shiota, Tatsuki; Kataoka, Kenichiro; Imai,
INVENTOR(S):
                         Minoru; Tsutsumi, Takaharu; Sudoh, Masaki;
                         Sogawa, Ryo; Morita, Takuya; Hada, Takahiko;
                         Muroga, Yumiko; Takenouchi, Osami; Furuya,
                         Monoru; Endo, Noriaki; Tarby, Christine M.;
                         Moree, Wil A.; Teig, Steven L.
                         Teijin Ltd., Japan; Combichem, Inc.
PATENT ASSIGNEE(S):
                         PCT Int. Appl., 374 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
                         English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
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PATENT INFORMATION:
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PATENT NO. KIND DATE APPLICATION NO. DATE																
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AU	9913	741		A.	1	1999	0607		Α	U 19	99-1	3741		1998	1117	
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US 1998-55285 A 19980406 US 1998-133434 A 19980813 WO 1998-US23254 W 19981117

OTHER SOURCE(S):

MARPAT 131:18925

GI

$$\begin{array}{c|c}
R^{1} & O & R^{4} \\
R^{2} & CH_{2} & N & CH_{2} & N & CH_{2} & N & CH_{2} & N & N \\
R^{2} & R^{3} & R^{3} & R^{4} & CH_{2} & R^{4} & CH_{2} & R^{6}
\end{array}$$

The title compds. [I; R1 = (un) substituted Ph, cycloalkyl, AB heteroaryl, etc.; R2 = H, alkyl, alkoxycarbonyl, etc.; j = 0-2; k =0-2; m = 2-4; n = 0-1; R3 = H, alkyl; R4, R5 = H, OH< Ph, etc.; p = 0-1; q = 0-1; G = CO, SO, CO2, etc.; R6 = Ph, cycloalkyl, cycloalkenyl, etc.] and their pharmaceutically acceptable acid addn. salts which inhibit the action of chemokines such as MIP-1.alpha. and/or MCP-1 on target cells and may be useful as a therapeutic drug and/or preventative drug in diseases, such as atherosclerosis, rheumatoid arthritis, and the like where blood monocytes and lymphocytes infiltrate into tissues, were prepd. Thus, reaction of N-benzoylglycine with 3-amino-1-(4-chlorobenzyl)pyrrolidine.2HCl in the presence of 3-ethyl-1-[3-(dimethylaminopropyl)]carbodiimide. HCl, 1-hydroxybenzotriazole and Et3N in CHCl3 afforded 95% II which showed 50-80% inhibition of MIP-1.alpha. binding to THP-1 cells at 10 .mu.M.

IT 226241-50-5P 226241-52-7P 226241-63-0P 226241-64-1P 226241-65-2P 226241-66-3P 226241-67-4P 226241-68-5P 226241-69-6P 226241-82-3P 226241-83-4P 226242-54-2P 226243-23-8P 226243-25-0P 226243-27-2P 226243-29-4P 226245-19-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of cyclic amine derivs. for inhibition of the action of chemokines such as MIP-1.alpha. and/or MCP-1 on target cells)

REFERENCE COUNT:
6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L16 ANSWER 30 OF 43 CAPLUS COPYRIGHT 2002 ACS 1999:282200 CAPLUS ACCESSION NUMBER: 130:311817 DOCUMENT NUMBER: Preparation of piperidine and piperazine TITLE: glycoprotein IIb/IIIa antagonists Carceller, Elena; Jimenez, Pere J.; Salas, Jorge INVENTOR(S): J. Uriach & Cia. S.A., Spain PATENT ASSIGNEE(S): PCT Int. Appl., 97 pp. SOURCE: CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. DATE KIND DATE PATENT NO. ---------_____ WO 1998-EP6751 19981023 WO 9920606 A2 19990429 WO 9920606 A3 19990429 AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG 19990510 AU 1999-21513 19981023 AU 9921513 . A1 ES 1997-2188 19971023 PRIORITY APPLN. INFO.: WO 1998-EP6751 19981023 MARPAT 130:311817 OTHER SOURCE(S): R1Z(CH2)mZ1Z2R [I; R = CO2H or metabolically labile ester or amide group; R1 = 1 - (4-piperidinyl) - 4-piperidinyl, 4-(4-piperidinyl) - 1piperidinyl, 4-(4-piperidinyl)-1-piperazinyl, etc.; Z = phenylene, pyridinediyl, pyrimidinediyl, etc.; Z1 = CONH, NHCO, SO2NH, etc.; Z2 - (un)substituted alkylene; Z1 = CO and Z = 1,n-azacycloalkylene; m = 0 or 1] were prepd. Thus, N-protected 4-R1C6H4CO2H (R1 = 4,4'-bipiperidin-1-yl)(prepn. given) was amidated by H2NCH2CH2CO2Me to give, after deprotection and sapon., 4-R1C6H4CONHCH2CH2CO2H (R1 = 4,4'-bipiperidin-1-yl). Data for biol. activity of I were given. 223535-05-5P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (prepn. of piperidine and piperazine glycoprotein IIb/IIIa antagonists) 223535-91-9P TT RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (prepn. of piperidine and piperazine glycoprotein IIb/IIIa antagonists) L16 ANSWER 31 OF 43 CAPLUS COPYRIGHT 2002 ACS 1999:48694 CAPLUS ACCESSION NUMBER: 130:124898 DOCUMENT NUMBER:

Preparation of 2-(4-bromo or 4-iodo TITLE:

phenylamino) benzoic acid derivatives as MEK

inhibitors

Barrett, Stephen Douglas; Bridges, Alexander INVENTOR(S):

James; Cody, Donna Reynolds; Doherty, Annette Marian; Dudley, David Thomas; Saltiel, Alan Robert; Schroeder, Mel Conrad; Tecle, Haile

PATENT ASSIGNEE(S): Warner-Lambert Company, USA

PCT Int. Appl., 67 pp. SOURCE:

CODEN: PIXXD2 Patent

DOCUMENT TYPE: LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAC	PATENT NO.					DATE			A	PPLI	CATI	ON NO	ο.	DATE		
WO	9901	 421		 A	 1	1999	0114		W	0 19	98-U	S131	- - 05	1998	0624	
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•														AM,		
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	9882															
EP	9934	37		A.	1	2000	0419		E	P 19	98-9	3282	9	1998	0624	
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	9810													1998		
JP	2002	5095	36	T	2	2002	0326		J	P 19	99-5	0722	7 .			
ZA	9805	726		Α		1999	0127		Z	A 19	98-5	726		1998		
	6310													2000		
US	2002	0226	47	A.	1	2002	0221							2001		
PRIORITY	APP	LN.	INFO	.:								-		1997		
														1998		
OM!! T.D. 0/						ח מ כו				000-	4623	19	A3	2000	0105	

OTHER SOURCE(S):

MARPAT 130:124898

GI

AB The title compds. [I; R1 = H, OH, C1-8 alkyl, etc.; R2 = H; R3-R5 = H, OH, halo, etc.; Z = COOR7, tetrazolyl, CONR6R7, etc.; R6, R7 = H, C1-8 alkyl, C2-8 alkenyl, etc.], which are potent inhibitors of MEK and, as such, are effective in treating cancer and other proliferative diseases such as inflammation, psoriasis and restenosis, as well as stroke, heart failure, and immunodeficiency

> 308-4994 Searcher : Shears

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disorders, were prepd. and formulated. Thus,
    treatment of 2-amino-5-iodotoluene in THF with LDA in
    THF/heptane/ethylbenzene soln. followed by addn. of
    2,4-difluorobenzoic acid in THF afforded II which showed IC50 of
    0.019 .mu.M against MEK in vitro.
    212628-77-8P 212628-80-3P 212628-81-4P
TΤ
    212628-82-5P 212628-83-6P 212628-85-8P
    212628-86-9P 212628-87-0P 212628-88-1P
    212628-89-2P 212628-90-5P 212628-91-6P
    212628-92-7P 212628-93-8P 212628-94-9P
    212628-99-4P 212629-00-0P 212629-01-1P
    212629-02-2P 212629-03-3P 212629-04-4P
    212629-05-5P 212629-06-6P 212629-07-7P
    212629-08-8P 212629-09-9P 212629-10-2P
    212629-11-3P 212629-12-4P 212629-13-5P
    212629-14-6P 212629-15-7P 212629-16-8P
    212629-17-9P 212629-18-0P 212629-19-1P
    212629-20-4P 212629-21-5P 212629-22-6P
    212629-23-7P 212629-24-8P 212629-25-9P
    212629-26-0P 212629-27-1P 212629-28-2P
    212629-29-3P 212629-30-6P 212629-31-7P
    212629-32-8P 212629-33-9P 212629-34-0P
    212629-35-1P 212629-36-2P 212629-37-3P
    212629-38-4P 212629-39-5P 212629-40-8P
    212629-41-9P 212629-42-0P 212629-43-1P
    212629-44-2P 212629-46-4P 212629-47-5P
    212629-48-6P 212629-50-0P 212629-52-2P
    212629-54-4P 212629-56-6P 212629-58-8P
    212629-60-2P 212629-62-4P 212629-64-6P
    212629-66-8P 212629-68-0P 212629-69-1P
    212629-71-5P 212629-73-7P 212629-75-9P
    212629-77-1P 212629-78-2P 212629-79-3P
    212629-80-6P 212629-81-7P 212629-82-8P
    212629-83-9P 212629-84-0P 212629-85-1P
    212629-86-2P 212629-87-3P 212629-88-4P
    212629-89-5P 212629-90-8P 212629-91-9P
    212629-92-0P 212629-93-1P 212630-00-7P
    212630-03-0P 212630-06-3P 212630-07-4P
    212630-08-5P 212630-09-6P 212630-10-9P
    212630-11-0P 212630-12-1P 212630-14-3P
    212630-15-4P 212630-16-5P 212630-17-6P
    212630-18-7P 212630-19-8P 212630-20-1P
    212630-21-2P 212630-22-3P 212630-23-4P
    212630-24-5P 212630-25-6P 212630-27-8P
    212630-28-9P 212630-29-0P 212630-30-3P
    212630-31-4P 212630-32-5P 212630-33-6P
     212630-34-7P 212630-35-8P 212630-36-9P
     212630-37-0P
    RL: BAC (Biological activity or effector, except adverse); BSU
     (Biological study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
     (Uses)
        (prepn. of 2-(4-bromo or 4-iodo phenylamino)benzoic acid derivs.
        as MEK inhibitors)
                               THERE ARE 17 CITED REFERENCES AVAILABLE
REFERENCE COUNT:
                         17
                               FOR THIS RECORD. ALL CITATIONS AVAILABLE
                               IN THE RE FORMAT
```

L16 ANSWER 32 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1998:785657 CAPLUS

DOCUMENT NUMBER:

130:38644

TITLE:

Preparation of ring-expanded nucleosides and

nucleotides as virucides and bactericides

INVENTOR(S):

Hosmane, Ramachandra; Burns, Barry

PATENT ASSIGNEE(S): SOURCE:

Universy of Maryland, USA; Nabi U.S., 24 pp., Cont.-in-part of U.S. Ser. No.

268,570, abandoned.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5843912	A	19981201	US 1995-518278	19950823
PRIORITY APPLN. INFO.	:	US	1994-268570	19940706
OTHER SOURCE(S):	MA	RPAT 130:38644		

OTHER SOURCE(S):

GI

AB The present invention relates to compns. comprising analogs of purine nucleosides contg. a ring-expanded ("fat") heterocyclic ring, I (R1, R3, R5 = independently NH, NH2, O, OH, S, SH. NH-alkyl, N-alkyl, O-alkyl, S-alkyl, NH-aryl, O-aryl, S-aryl; R2, R4, R7, R8 = independently , H, alkyl, substituted Ph, heterocycle, aralkyl; R6 = H, alkyl, Ph, substituted Ph, heterocycle, aralkyl, glycosyl, ; U, X, Y, Z, W, J, K, L = C, N) in place of purine, and an unmodified or modified sugar residue, pharmaceutically acceptable derivs. of such compns., as well as methods of use thereof. In particular, these compns. may be utilized in the treatment of certain cancers, bacterial, fungal, parasitic, and viral infections, including, but not limited to, Acquired Immunodeficiency Syndrome (AIDS) and hepatitis. 6-Amino-6-methoxycarbonyl-4,5,7,8-tetrahydro-6H-imidazo[4,5,e]-[1,4]-diazepine-5,8-dione was prepd. as adenosine deaminase and guanase inhibitor and tested for its anti-retroviral and antibacterial activities.

IT 169317-90-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

> Shears 308-4994 Searcher :

(prepn. of ring-expanded nucleosides and as virucides and

bactericides)

THERE ARE 1 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT: 1

THIS RECORD. ALL CITATIONS AVAILABLE IN

THE RE FORMAT

L16 ANSWER 33 OF 43 CAPLUS COPYRIGHT 2002 ACS

1998:236274 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 128:282780

Preparation of heterocyclic inhibitors of TITLE:

microsomal triglyceride transfer protein

Biller, Scott A.; Dickson, John K.; Lawrence, R. INVENTOR(S):

Michael; Magnin, David R.; Poss, Michael A.;

Sulsky, Richard B.; Tino, Joseph A.

Bristol-Myers Squibb Co., USA PATENT ASSIGNEE(S):

U.S., 185 pp. Cont.-in-part of U.S. Ser. No. SOURCE:

391,901, abandoned.

CODEN: USXXAM

DOCUMENT TYPE:

Patent English

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5739135 CA 2091102 HU 67962	A AA	19980414 19930907 19950529	US 1995-472067 CA 1993-2091102 HU 1993-627	19950606
HU 218419 JP 06038761 EP 584446 EP 584446	A2 A2 A3		JP 1993-46499 EP 1993-103697	
R: AT, BE, PT, SE	CH, DE,		GB, GR, IE, IT, LI,	LU, MC, NL,
AU 670930 AU 9334064	B2 A1	19930909	AU 1993-34064	
US 5595872 US 5789197	A	19980804	US 1993-117362 US 1995-486924	
US 5712279 IL 116917	A Al	19980127 20000831	US 1996-548811 IL 1996-116917 CA 1996-2213466	19960111
WO 9626205	AA A1	19960829	WO 1996-US824	19960201
W: AU, BG,	CA, CN,	CZ, EE, FI, SG, SK, UA	GE, HU, JP, KR, LT,	LV, MX, NO,
RW: AT, BE, SE	CH, DE,	DK, ES, FR,	GB, GR, IE, IT, LU,	
AU 699865	B2	19981217	AU 1996-47631	
	A1	19981230	CN 1996-192015 EP 1996-903604	19960201
PT, IE		•	GB, GR, IT, LI, LU,	
ZA 9601340	T2 A	19970911	JP 1996-525679 ZA 1996-1340	19960201 19960220
US 5883099 US 6034098	Α	20000307	US 1997-896872 US 1997-898304	
US 6066650 FI 9703416	A A		US 1997-898303 FI 1997-3416	19970721

308-4994 Searcher : Shears

19970820 NO 1997-3821 19970820 NO 9703821 Α LT 1997-152 LT 4367 в 19980825 19970919 LV 1997-171 В 19981120 19970919 LV 11951 US 1993-117362 A2 19930903 PRIORITY APPLN. INFO.: US 1994-284808 B2 19940805 US 1995-391901 B2 19950221 US 1992-847503 A 19920306 US 1993-15449 B2 19930222 US 1995-472067 A2 19950606 WO 1996-US824 W 19960201

OTHER SOURCE(S):

MARPAT 128:282780

GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The title compds. [I-V; Q = C(O), S(O)2; X = CHR8, C(O), CHR9CHR10, AB CR9:CR10 (wherein R8-R10 = H, alkyl, alkenyl, etc.); Y = (CH2)m, C(0) (m = 2-3); R1 = alkyl, alkenyl, alkynyl, etc.; R2-R4 = H, halo, alkyl, etc.; R5 = alkyl, alkenyl, alkynyl, etc.; R6 = H, C1-4 alkyl, C1-4 alkenyl] which inhibit microsomal triglyceride transfer protein and thus are useful for lowering serum lipids and treating atherosclerosis and related diseases such as hyperglycemia and obesity, were prepd. Thus, reaction of 1-(3,3-diphenylpropyl)-4piperidinamine. HCl (prepn. described) with benzoyl chloride in the presence of Et3N in CH2Cl2 afforded 84% the title compd. III.HCl [Q = C(0); R1 = 3,3-diphenylpropyl; R5 = Ph; R6 = H]. Compds. I-V are effective at 5-500 mg/day.

163267-27-4P 182429-76-1P 182429-79-4P ITRL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of heterocyclic inhibitors of microsomal triglyceride transfer protein)

L16 ANSWER 34 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

1998:115356 CAPLUS 128:154011

TITLE:

Preparation of 9-thioxanthenecarboxamides and

9-fluorenecarboxamides as inhibitors of microsomal triglyceride transfer protein

INVENTOR(S):

Biller, Scott A.; Dickson, John K.; Lawrence, R.

Michael; Magnin, David R.; Poss, Michael A.; Robl, Jeffrey A.; Sulsky, Richard B.; Tino,

Joseph A.

PATENT ASSIGNEE(S):

Bristol-Myers Squibb Co., USA

SOURCE:

U.S., 98 pp., Cont.-in-part of U.S. Ser.

No.472,067. CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

	5712 2091			A A		1998 1993						96-5 93-2		_		1996 1993		
	6796			A2		1995						93-			_	1993		
HU	2184	19		В		2000	0828											
. JP	0603	8761		A2	2	1994	0215			JΡ	19	93-4	464	99		1993	0308	
EP	5844	46		A2	2	1994	0302			ΕP	19	93-3	103	697	7	1993	0308	
EP	5844	46		A.		1995												
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB	, (GR,	IE,	, I	Τ,	LI,	LU,	MC,	NL,
		PT,	SE															
AU	6709	30		B	2	1996	8080			ΑU	19	93-3	340	64		1993	0309	
AU	9334	064		A.	L	1993	0909											
US	5739	135		Α		1998	0414			US	19	95-4	472	067	7	1995	0606	
ZA	9601	340		Α		1997	0911			ZA	19	96-:	134	0		1996	0220	
LT	4367			В		1998	0825			LT	19	97-:	152			1997	0919	
PRIORITY	APP	LN.	INFO.	. :				1	US	199	95-3	391	901		В2	1995	0221	
								1	US	199	95-	4720	067		Α2	1995	0606	
								1	US	199	92-	347	503		Α	1992	0306	
								1	US	199	93-:	1173	362		A2	1993	0903	
								1	US	199	94-	2848	308		В2	1994	0805	

OTHER SOURCE(S): MARPAT 128:154011

GI

$$X^{1}$$
 $C(0) NHCH_{2}CF_{3}$
 Z
 $NHC(0) R^{5}$
 X^{2}
 I

The title compds. [I; Z = a bond, S; X1, X2 = H, halo; x = 2-6; (CH2)x is optionally substituted with 1-3 substituents such as alkyl or halo; R5 = (un)substituted heteroaryl, aryl, heterocycloalkyl, cycloalkyl] and their piperidine N-oxides, which inhibit microsomal triglyceride transfer protein and thus are useful for preventing or

treating atherosclerosis, pancreatitis secondary to hypertriglyceridemia, hyperglycemia, or obesity, and for lowering serum lipid levels, or preventing and/or treating hyperlipemia, hyperlipidemia, hyperlipoproteinemia, hypercholesterolemia, and/or hypertriglyceridemia, were prepd. Thus, reaction of 9-fluorenecarboxamide II (prepn. of both reagents is described) with piperidine III in PhMe/DMF afforded the title compd. I [Z = a bond; X1 = X2 = H; (CH2)x = (CH2)2CF2CH2; R5 = 2-biphenyl]. Compds. I are effective at 5-500 mg/day.

IT 182431-88-5P 182432-11-7P 182434-95-3P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of 9-thioxanthenecarboxamides and 9-fluorenecarboxamides as inhibitors of microsomal triglyceride transfer protein)

L16 ANSWER 35 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1995:858623 CAPLUS

DOCUMENT NUMBER: 123:256357

TITLE: Preparation of anthranilic acid amide derivative

as cyclic guanosine monophosphate-

phosphodiesterase inhibitors

INVENTOR(S): Ozaki, Fumihiro; Ishibashi, Keiji; Ikuta,

Hironori; Ishihara, Hiroki; Souda, Shigeru

PATENT ASSIGNEE(S): Japan

SOURCE: PCT Int. Appl., 204 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO. DATE
	A1 19950706	WO 1994-JP2262 19941227
W: AU, CA,	CN, FI, HU, KR, NO,	NZ, RU, US
RW: AT, BE,	CH, DE, DK, ES, FR,	GB, GR, IE, IT, LU, MC, NL, PT,
SE		
CA 2155662	AA 19950706	CA 1994-2155662 19941227
AU 9512824	A1 19950717	AU 1995-12824 19941227
AU 694465	B2 19980723	
EP 686625	A1 19951213	EP 1995-903999 19941227
	B1 19990526	
R: AT, BE,	CH, DE, DK, ES, FR,	GB, GR, IE, IT, LI, LU, MC, NL,
PT, SE		
CN 1118595	A 19960313	CN 1994-191311 19941227
JP 08188563	A2 19960723	JP 1994-336920 19941227
HU 74450		HU 1995-2512 19941227
RU 2128644		RU 1995-120194 19941227
Δ Ͳ 180468	E 19990615	АТ 1995-903999 19941227
FT 9503968	A 19951019	FI 1995-3968 19950823
NO 9503305	A 19951025	NO 1995-3305 19950823
	A 19980210	
PRIORITY APPLN. INFO		JP 1993-347092 A 19931227
INIONIII MIIDM. IMIO		JP 1994-299110 A 19941109
		WO 1994-JP2262 W 19941227
OTHER SOURCE(S):		

GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Anthranilamide derivs. [I; R1, R2, R3, R4 = H, halo, OH, AB (halo)alkyl, (halo)alkoxy, nitro, hydroxyalkyl, cyano, (CH2)pNR9R10, S(O)qR13, (un)protected CO2H, (un)substituted tetrazolyl, CONH2, pyrazolyl, or imidazolyl; or adjacent two substituents selected from R1 - R4 together with the C atoms bonded to them forms a ring; wherein R9, R10 = H, (halo)alkyl, arylalkyl, heteroarylalkyl, acyl, (un)protected CO2H; or NR9R10 forms a ring; p = 0, 1-6; R13 = H, (halo)alkyl; q = 0, 1-2; R5, R6 = H, halo, OH, cyano, (halo)alkyl,(halo)alkoxy; or R5 and R6 together with the C atoms bonded to them form cycloalkane, oxolane, 1,3-dioxolane, or 1,4-dioxane ring; W = N, CH; R7, R8 = H, (halo)alkyl; or R1 and R7 together with the C atoms bonded to them form a ring optionally contg. other N, O, or S atom; A = H, (halo)alkyl, X(CH2)mZ; wherein X = CO, CS, CH2, SO2; Z= OH, (halo) alkoxy, cyano, halo, etc.; Y = O, S; n = 0, 1-6] or pharmacol. acceptable salts thereof are prepd. These compds. are useful for the treatment of ischemic heart disease , angina pectoris, hypertension, pulmonary hypertension, heart failure, and asthma. Thus, 2-nitro-5-chlorobenzoic acid was refluxed with SOC12 in benzene for 4 h and concd. to give 2-nitro-5-chlorobenzoyl chloride which was amidated with piperonylamine in the presence of Et3N in THF to give a benzamide This compd. was reduced by Fe powder in a mixt. of (II; R = NO2).AcOH, H2O, and MeOH under gentle refluxing to give, after concn. and treatment with concd. HCl in EtOH, N-piperonylanthranilamide deriv. II. HCl (R = NH2). An anthranilamide deriv. (III) showed IC50 of 0.4 nM against cyclic guanosine monophosphatephosphodiesterase prepn. from pig aorta.

IT 169043-59-8P

> RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of anthranilamide derivs. as cyclic guanosine monophosphate-phosphodiesterase inhibitors)

L16 ANSWER 36 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1995:568500 CAPLUS DOCUMENT NUMBER:

123:169516

TITLE:

Preparation of acylaminopiperidines and piperazines as inhibitors of microsomal

triglyceride transfer protein.

INVENTOR(S):

Wetterau, John R., II; Sharp, Daru Young; Gregg, Richard E.; Biller, Scott A.; Dickson, John K.; Lawrence, Michael R.; Lawson, John E.; Holava,

Henry M.; Partyka, Richard A. Bristol-Myers Squibb Co., USA

PATENT ASSIGNEE(S):

Eur. Pat. Appl., 134 pp.

SOURCE: CODEN: EPXXDW

DOCUMENT TYPE:

Patent English

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

308-4994 Shears Searcher :

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
EP 643057	A1 199503	5 EP 1994-113800	19940902
Dm Cr		, FR, GB, GR, IE, IT, LI,	
CA 2091102	AA 199309	7 CA 1993-2091102 5 ZA 1993-1601 9 HU 1993-627 8 5 JP 1993-46499 2 EP 1993-103697 6	19930305
ZA 9301601	A 199310	5 ZA 1993-1601	19930305
ни 67962	A2 199505	9 ни 1993-627	19930305
HU 218419	B 2000083	8	
JP 06038761	A2 199402	5 JP 1993-46499	19930308
EP 584446	A2 199403	2 EP 1993-103697	19930308
EP 584446	A3 1995042	6	
R: AT, BE	, CH, DE, DK, ES	, FR, GB, GR, IE, IT, LI,	LU, MC, NL,
PT. SE			
AU 670930	B2 1996080 A1 1993090	8 AU 1993-34064	19930309
AU 9334064	A1 1993090	9	
US 5595872 CA 2131430 FI 9404048	A 1997012	1 US 1993-117362	19930903
CA 2131430	AA 1995030		19940902
FI 9404048	A 1995030	4 FI 1994-4048	19940902
NO 9403260	A 1995030	6 NO 1994-3260	19940902
AU 9471642	A1 199503	6 AU 1994-71642	19940902
AU 690125	B2 199804:	NO 1994-3260 AU 1994-71642	
ZA 9406772	A 1995040	3 ZA 1994-6//2	19940902
JP 07165712	A2 1995062	7 JP 1994-210057	19940902
CN 1106003	A 1995080	2 CN 1994-115640	19940902
ни 70613	A2 199510	7 JP 1994-210057 2 CN 1994-115640 0 HU 1994-2542	19940902
US 5789197	A 1998080	4 US 1995-486924	19950607
RIORITY APPLN. IN	0.:	US 1993-117362 A	
		US 1992-847503 A	
		US 1993-15449 B2	19930222
THER SOURCE(S):	MARPAT 12	:169516	

$$R^{2}$$
 R^{3}
 R^{4}
 NR^{1}
 $R^{5}CON_{R}^{6}$
 NR^{1}
 R^{2}
 R^{3}
 R^{2}
 R^{3}
 R^{4}
 NR^{7}
 R^{6}
 R^{7}
 R^{7}

Title compds. [I-III; X = CHR8, CHR9CHR10, CR9:CR10; R8-R10 = H, AB alkyl, alkenyl, alkynyl, aryl, aralkyl, heteroaryl, heteroarylalkyl, cycloalkyl, cycloalkylalkyl; Y = (CH2)m, CO; m = 2, 3; R1 = (substituted) alkyl, alkenyl, alkynyl, aryl, heteroaryl, aralkyl, diarylalkyl, diarylalkenyl, diarylalkynyl, diarylalkylaryl, heteroarylalkyl, cycloalkyl, cycloalkylalkyl, etc.; R2-R4 = H, halo, alkyl, haloalkyl, alkenyl, alkoxy, aryloxy, aryl, arylalkyl, alkylthio, arylthio, cycloalkyl, cycloalkylalkyl, heteroaryl, heteroarylalkyl, OH, haloalkyl; R5 = (substituted) alkyl, alkenyl, alkynyl, aryl, heteroaryl, aralkyl, heteroarylalkyl, cycloalkyl, cycloalkylalkyl, polycycloalkyl, cycloalkenyl, cycloalkenylalkyl, heteroarylcarbonyl, etc.; R6 = H, alkyl, alkenyl; R7 = alkyl, aryl, aralkyl, oxoalkyl, aryloxoalkyl], were prepd. as inhibitors of microsomal triglyceride transfer protein. Thus, tert-Bu 4-piperidinylcarbamate (prepn. given) and 3,3-diphenyl-1-propanol tosylate (prepn. given) were stirred with K2CO3 in Me2CHOH overnight to give 76% tert-Bu [1-(3,3-diphenylpropyl)-4-piperidinyl]carbamate. This was deprotected with 4N HCl in dioxane and the product was treated with PhCOCl and Et3N in CH2Cl2 to give title compd. (IV).

IT 163267-27-4P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of acylaminopiperidines and piperazines as inhibitors of microsomal triglyceride transfer protein)

L16 ANSWER 37 OF 43 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1995:242543 CAPLUS

DOCUMENT NUMBER:

122:31131

TITLE:

Preparation of benzamides in gastro-intestinal

pathologies.

Baldazzi, Claudia; Piani, Silvano; Barbanti, INVENTOR(S):

Maria; Marchi, Egidio

PATENT ASSIGNEE(S): Alfa Wassermann S.p.A., Italy

Eur. Pat. Appl., 20 pp. SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

GI

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	ENT 1	NO.		KIN	1D	DATE	3		AP	PLIC	CATIO	ON NO	ο.	DATE		
EP	6202	10		A1	L	1994	1019		EP	199	94-10	05463	3	19940	0408	
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	LI,				SE
CA	2120	214		AA	Ą	1994	1017		CA	. 199	94-23	1202		19940		
JP	0632	1881		A2	2	1994	1122					3637		19940	0412	
PRIORITY	APP	LN.]	INFO.	:					IT 19	93-E	30154	1		19930	0416	
OTHER SC	URCE	(S):			MAF	RPAT	122:3	3113	1							
O.T.																

$$R^{1}$$
 CONH (CH₂) mNR¹R²
 R^{6} NR³R⁴

Title compds. I (m = 1-4; R1, R2 = H, C1-6 alkyl, R1R2N = AB heterocyclyl; R3, R4 = H, C1-10 alkyl, PhCH2; R5-8 = H, C1-6 alkyl, halo) and salts thereof, are prepd. I have prokinetic effects, such as stimulation on gastro-intestinal motility, and possess anti-emetic qualities, without side effects involving the central nervous system. To 5-chloroisatoic anhydride in dimethylacetamide was added NaH to give 5-chloroN-methylisatoic anhydride to which in dioxane was added N, N-diethylaminoethylamine to give I (m = 2, R1 = R2 = Et, R3 = R5-7 - H, R4 = Me, R8 = C1) converted to the citrate. The biol. activity of I was demonstrated both in vitro and in vivo. IT 159619-35-9P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of benzamides in gastro-intestinal pathologies.)

L16 ANSWER 38 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1994:164213 CAPLUS

DOCUMENT NUMBER:

120:164213

Ι

TITLE:

Pyrido[2,3-d]pyrimidinone phosphodiesterase

inhibitors

INVENTOR(S):

Wilhelm, Robert Stephen; Chin, Ronnie Lipp;

Devens, Bruce Henry; Alvarez, Robert

PATENT ASSIGNEE(S):

Syntex (U.S.A.), Inc., USA

SOURCE:

PCT Int. Appl., 59 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

ם מונו

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	CENT N	10.		KI	ND	DAT	E		A	PPL	ICATI	ON N	ο.	DATE		
WO	93190 W:	ΑU,	CA,	FΙ,	HU,	JP	KR,	NO;	NZ							
		SE				DK	, ES,	FR,	GB,	GR	, IE,	IT,	LU	MC,	NL,	PT,
US	52644	37		Α		1993	31123		U	S 1	992-8	5517	9	1992	U33U	
AU	93391	86		A.	l	1993	31021		A	U 1	993-3	9186	•	19930		
AU	66952	0		B2	2	1996	50.613									
ZA	93019	45		Α		1994	10918		Z	A 1	993-1	945		1993	า จ.1 ผ	
EP	63158	0		A1	L	1995	0104		E	P 1	993-9	08322	2	19930	7318	
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR.	GB.	GR	, IE,	TT.	- Т.Т.	T.II	MC	NT.
		PT,	SE		•	•	•	•			,,	,	,	шо,	nc,	иш,
	67552				?	1995	0428		Н	J 1	994-2	653		19930	1318	
JP	07504	676		Т2	:	1995	0525				993-5			19930		
	32413					2001	.1225						-		,010	
IL	10509	2		A1		1998	0615		I	. 19	993-1	05092	2	19930	1318	
CN	10403	27		В		1998	1021		C	J 1	993-1	0335 2)	19930		
FI	94043	05		Α		1994	0916		F]	19	994-4	305		19940		
· NO	94034	56		Α		1994	0916		NO	19	994-3	456		19940		
PRIORITY	APPL	N. I	NFO.	:				U	S 19	92-	-8551	79	А	19920		
								W	0 19		-US22			19930		
OTHER SO	URCE (S):			MAR	PAT	120:1	6421	3			-	- -			

Ι

The title compds. I [R1 = H, (CH2)nR7; R7 = aryl, heteroaryl; n = 1, 2; R2-R6 = H, lower alkyl, halogen, CO2H, CO2Me, carbamoyl, etc.; Y = CH2, CO; only one of R2-R6 may be other than H], useful for the treatment of asthma, pain, inflammatory diseases, etc., are prepd. and I-contg. formulations presented. Thus, I (R1= 3-pyridylmethyl, R2 = R4 = R6 = H, R3 = NO2, Y = CO) was prepd. and demonstrated 50% inhibitory concn. for human lymphocyte cAMP phosphodiesterase (PDE 4) of 0.00026 .mu.M.

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (
(prepn. and reaction of, in prepn. of phosphodiesteral inhibitors)

L16 ANSWER 39 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

1993:625951 CAPLUS

119:225951

TITLE:

Preparation of cyclohexane- and

tetrahydro(thio)pyranthioformamide derivatives and analogs as ATP-sensitive potassium channel

openers

INVENTOR(S):

Kabasawa, Yasuhiro; Ozaki, Fumihiro; Ishibashi,

Keiji; Hasegawa, Takashi; Oinuma, Hitoshi; Ogawa, Toshiaki; Adachi, Hideyuki; Kato,

Hiroshi; Kodama, Kotaro; et al.

PATENT ASSIGNEE(S):

Eisai Co., Ltd., Japan PCT Int. Appl., 95 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese 1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATE	PATENT NO.			DATE		APPLICATION NO. DATE
				19930429 KR, NO,		WO 1992-JP1297 19921006
Ī	RW: AT,	BE,	CH, DE,	DK, ES,	FR,	GB, GR, IE, IT, LU, MC, NL, SE
JP 0 AU 9:	6025154 226742		A2 A1	19940201		JP 1992-265948 19921005 AU 1992-26742 19921006
AU 6	61044		B2	19950713		EP 1992-920995 19921006
						GB, GR, IE, IT, LI, LU, NL, SE
HU 6	6229		A2	19941028		HU 1994-1074 19921006 RU 1994-20984 19921006
KR 9	711277		В1	19970709		KR 1994-71154 19940409
						NO 1994-1294 19940411 FI 1994-1681 19940412
US 5	444066		A	19950822		US 1994-211701 19940426
US 5	498634		A	19960312		US 1995-380589 19950130 US 1995-531335 19950920
PRIORITY				19970223		JP 1991-264622 A 19911014
						JP 1992-197 A 19920106 WO 1992-JP1297 A 19921006
					1	US 1994-211701 A3 19940426
					Ţ	US 1995-380589 A3 19950130

OTHER SOURCE(S):

GI

MARPAT 119:225951

The title compds. [I; Y = 0, S(0)n (wherein n = 0-2), CO, CS, AB (un) substituted C(:CH2), C(:NH), CH2; Z = O, S(O)m (wherein m = CH2)

0-2), (CH2)p (wherein p = 0-2); A = (un)substituted aryl, thienyl, furyl, benzofurazanyl, pyrrolyl, imidazolyl, triazolyl, tetrazolyl, pyridinyl, pyrazolyl, isoxazolyl, isothiazolyl, oxazolyl, benzimidazolyl, imidazopyridyl, imidazopyrazinyl, imidazopyrimidinyl, etc., provided that when Y or Z = O or S A .noteq. unsubstituted imidazolyl; R1, R2 = H, lower alkyl, (un) substituted arylalkyl or heteroarylalkyl, or R1 R2 forms a benzene ring; R3, R4 = H, lower alkyl, cycloalkyl, lower alkoxy, OH, aryl, arylalkyl, heteroaryl, heteroarylalkyl, or R3R4 forms a ring optionally contg. O, N, or S], useful as antihypertensives and antiasthmatics and for the treatment of angina pectoris, are prepd. Thus, Grignard reaction of 6-bromoimidazo[1,2-a]pyridine with EtMqBr in refluxing THF followed by addn. reaction with 2-methoxycyclohexanone and hydrolysis with concd. H2SO4 to give 2-(imidazo[1,2-a]pyridin-6-yl)cyclohexanone (II). Stirring II with KOCMe3 in THF followed by addn. reaction with MeNCS in THF-DMF gave a imidazo[1,2-a]pyridinylcyclohexanecarbothiamide deriv. [(.+-.)-III; R = Me] which was resolved by (+)-dibenzoyl-D-tartaric acid monohydrate to give(-)-III (R = Me) (IV). IV and (-)-III (R = Me) Et) showed -log(IC50) of 5.58 and 6.16, resp., for shortening the action potential duration time (APD90) in isolated cardiac papillary muscles of guinea pigs and at 1 mg/kg p.o. reduced 22.1 and 40.9%, resp., the blood pressure of spontaneously hypertensive rats (SHR). A total of 25 I were prepd.

IT 150780-12-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as ATP-sensitive potassium channel opener)

L16 ANSWER 40 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1991:559174 CAPLUS

DOCUMENT NUMBER:

115:159174

TITLE:

Preparation of quinazoline-3-alkanoates as

platelet aggregation and aldose reductase

inhibitors

INVENTOR(S):

Fujimori, Shizuyoshi; Ohnota, Michiro; Hirata,

Yoshihiro; Murakami, Koji

PATENT ASSIGNEE(S):

Kyorin Pharmaceutical Co., Ltd., Japan

SOURCE:

PCT Int. Appl., 54 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent Japanese

LANGUAGE:

י. 1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	TENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO	9109024	A1	19910627	WO 1990-JP1600	19901210
	•	CA, HU, KR CH. DE. ES	, US , FR, GB, IT	r, NL, SE	
	03181469	A2	19910807	JP 1989-321097	19891211
JΡ	07047582	B4	19950524		
ÇA	2046603	AA	19910612	CA 1990-2046603	19901210
ΑU	9168905	A1	19910718	AU 1991-68905	19901210
ΑU	640194	B2	19930819		
ΕP	456835	A1	19911121	EP 1991-900052	19901210
EΡ	456835	B1	19960515		
	R: BE, (CH, DE, ES	, FR, GB, IT	r, LI, NL, SE	
HU	58304	A2	19920228	ни 1991-2399	19901210

HU 207999 B 19930728 ES 2087991 T3 19960801 ES 1991-900052 19901210 US 5234928 A 19930810 US 1991-721610 19910717

PRIORITY APPLN. INFO.: JP 1989-321097 19891211 WO 1990-JP1600 19901210

OTHER SOURCE(S): MARPAT 115:159174

Ι

GI

$$\begin{array}{c|c}
R^2 & X & N(CH_2)_nCO_2R \\
N & O \\
R^3 & Ar^1
\end{array}$$

The title compds. [I; R = H, carboxy-protective group; R1 = alkyl, AB alkenyl, alkynyl, alkoxy, alkylthio, halo, (substituted) Ph, heterocyclyl, or benzoyl, naphthyl, cycloalkyl; R2, R3 = H, halo, alkyl, alkoxy, (substituted) aralkyl, NO2, imidazolyl, imidazolylmethyl, NR4R5; R4, R5 = H, alkyl; or NR4R5 = 5- or 6-membered heterocyclyl optionally contg. other heteroatom(s); X =CO, C(S), (alkyl-substituted) CH2; A = alkylene, alkenylene; n = alkylene1-3], useful for treatment of thrombosis, heart diseases, or diabetes complications, are prepd. Thus, condensation of H2NCH2CO2Et.HCl with 6-chloro-2H-3,1-benzoxazine- $2,4\,(\mbox{1H})\mbox{-dione}$ in dioxane contg. Et3N and cyclocondensation of the resulting 2,5-(H2N)ClC6H3CONHCH2CO2Et with N,N'-carbonyldiimidazole in dioxane at 150.degree. gave Et 6-chloro-1,4-dihydro-2,4-dioxo-3(2H)-quinazolineacetate which was alkylated with 2-ClC6H4CH2Cl in the presence of NaH in DMF at 70.degree. to give Et 6-chloro-1-(4-chlorophenyl)methyl)-1,4-dihydro-2,4-dioxo-3(2H)quinazolineacetate. A total of 196 I were prepd. and in vitro inhibited aldose reductase with IC50 of 10-7 - 10-8 M and arachidonic acid-induced rabbit's platelet aggregation with IC50 of 10-5 - 10-7 M.

IT 136148-82-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of, as intermediate for aldose reductase and platelet aggregation inhibitor quinazolinealkanoic acid deriv.)

IT 136148-82-8

RL: RCT (Reactant)

(reaction of, in prepn. of aldose reductase and platelet aggregation inhibitor quinazolinealkanoic acid deriv.)

L16 ANSWER 41 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1991:207038 CAPLUS

DOCUMENT NUMBER:

114:207038

TITLE:

Preparation of 1-pyridyl-2-(substituted

amino) cyclohexanecarbothioamides and analogs as

smooth muscle relaxants

INVENTOR(S):

Hart, Terance William; Vacher, Bernard Yvon

Jack; Walsh, Roger John Aitchison

PATENT ASSIGNEE(S):

Rhone-Poulenc Sante, Fr.

SOURCE:

Eur. Pat. Appl., 18 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent English

LANGUAGE:

NGUAGE: E

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 403398	A1	19901219	EP 1990-401735	19900615
EP 403398 R: AT, BE,				•
AU 9057128 ZA 9004621	Al A	19901220 19910327	ZA 1990-4621	19900614 19900614
CA 2019106 JP 03063260	AA A2	19901216 19910319	JP 1990-157400	19900615 19900615
HU 54983 ES 2064681	A2 T3	19910429 19950201		19900615 19900615
US 5276045 PRIORITY APPLN. INFO	A	19940104	00 2002 00000	19920330 19890616
				19890616 19900615

OTHER SOURCE(S):

MARPAT 114:207038

GΙ

The title compds. [I; A = (un)substituted N-contg. heterocyclyl, Ph; R = alkyl; when n = 0, Rl = H, acyl, (un)substituted (cyclo)alkyl, aryl, etc.; when n = 1, Rl = (un)substituted alkyl, PhCH2, naphthylmethyl, pyridylmethyl, etc.; R2 = groups cited for Rl (n = 0); n = 0, 1; m = 0-2] were prepd. for prophylaxis and/or treatment of disorders assocd. with vascular, respiratory, or gastrointestinal smooth muscle contraction. Thus, 2-(3-pyridyl)cyclohexanone was condensed with (R)-PhMeCHNH2 and the product treated, sequentially, with BuLi and MeNCS in THF to give pyridylcyclohexanecarbothioamide II (R2R3 = bond) which was reduced with NaBH3CN to give (2R,1S)-II (R2 = R3 = H) which had EC90 of 10-5 .mu.M for redn. of K+-induced contractions of rat aorta strips.

IT 133667-58-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, in prepn. of smooth muscle relaxants)

IT 133667-59-1P 133670-75-4P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (prepn. of, as smooth muscle relaxant)

L16 ANSWER 42 OF 43 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1987:156252 CAPLUS

DOCUMENT NUMBER:

106:156252

TITLE:

SOURCE:

AUTHOR(S):

CORPORATE SOURCE:

Potential antitumor agents. 50. In vivo

solid-tumor activity of derivatives of

N-[2-(dimethylamino)ethyl]acridine-4-carboxamide

Atwell, Graham J.; Rewcastle, Gordon W.;

Baguley, Bruce C.; Denny, William A.

Sch. Med., Univ. Auckland, Auckland, N. Z.

J. Med. Chem. (1987), 30(4), 664-9

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 106:156252

GI

The synthesis, physicochem. properties, and antitumor activity of a AB series of N-[2-(dialkylamino)alkyl]acridine-4-carboxamides (e.g., I; R = H, 5-Cl, 8-Me) are reported. Thus, the K salt of oxoacridancarboxylic acid II was treated with Al amalgam in aq. EtOH, followed by treatment with HCl and FeCl3, to give 64% the acridinecarboxylic acid III, which was treated with 1,1'-carbonyldiimidazole in DMF and then Me2NCH2CH2NH2 to give 60% I (R = H). The title compds. bind to DNA by intercalation, but exist under physiol. conditions as monocations due to the weakly basic acridine chromophore (pKa = 3.5-4.5). The acridine-4-carboxamides show very broad structure-activity relationships (SAR) for antileukemic activity, with substituents at nearly all acridine positions proving acceptable. The compds. also show remarkable activity against the Lewis lung solid tumor in vivo, with several analogs (e.g., I; R = H) capable of effecting 100% cures of the advanced disease. The broad SAR and high solid-tumor activity of the 9-acridine-4-carboxamides imply they should be considered as a completely new class of antitumor agent. IΤ 89459-32-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and intramol. cyclocondensation of)

L16 ANSWER 43 OF 43 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1979:592973 CAPLUS

DOCUMENT NUMBER:

91:192973

2-(Substituted amino)ethanol nitrate esters TITLE: Nagano, Hiroyuki; Matsunaga, Isao; Shindo, INVENTOR(S):

Minoru

PATENT ASSIGNEE(S):

Chugai Pharmaceutical Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE -----_____ JP 54081222 19790628 JP 1977-147109 19771209 A2

GI

Nineteen nitrate esters RR1NCH2CH2ONO2 (I; RR1N = AB 3,4,5-trimethoxybenzamido, o-AcOC6H4CONH, o-EtO2CC6H4O2CNH, 3-pyridinesulfonamido, 2,3-pyridinedicarboximido, etc.), e.g., II, III.HCl (R2 = 2-methoxyphenyl, 3,4,5-trimethoxyphenyl, 3-pyridyl), or IV.HCl, useful for treating circulatory disorders (no data), were prepd. by, e.g., acylating H2NCH2CH2ONO2 (V). IV.HCl was prepd. by NaBH4 redn. of a Schiff base from pyridoxal and V. Thus, 20 mL C6H6 satd.with COC12 was treated dropwise with 1 g V and 1 g Et3N in Et2O, excess COC12 was removed, and the mixt. was stirred with 2.47 g methoxamine hydrochloride and aq. NaHCO3 in EtOAc to give 0.5 g II.

71908-18-4P 71908-19-5P ΤΤ

> RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

=> sel hit 116 1-43 rn E1 THROUGH E314 ASSIGNED

REGISTRY ' ENTERED AT 14:48:45 ON 31 MAY 2002 L17 314 SEA FILE=REGISTRY ABB=ON PLU=ON (226241-69-6/BI OR 212628-77-8/BI OR 212628-80-3/BI OR 212628-81-4/BI OR 212628-82-5/BI OR 212628-83-6/BI OR 212628-85-8/BI OR 212628-86-9/BI OR 212628-87-0/BI OR 212628-88-1/BI OR

> Shears 308-4994 Searcher :

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212630-03-0/BI OR 212630-06-3/BI OR 212
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1,2,5-7,14-16,23,51,52,54,55,57,58,63,66-71,73-76,79,81,82,84,91-94,130,150,151,156,155,167,169,201,280,296-310,312,313 ide can

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L17 ANSWER 1 OF 314 REGISTRY COPYRIGHT 2002 ACS
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RN 408369-40-4 REGISTRY

CN Benzamide, N-[3-fluoro-4-[[2-(2-pyridinyl)ethyl]amino]phenyl]-2-[[3-(trifluoromethyl)phenyl]amino]- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN N-[3-Fluoro-4-[[2-(2-pyridinyl)ethyl]amino]phenyl]-2-[3-(trifluoromethyl)anilino]benzamide

FS 3D CONCORD

MF C27 H22 F4 N4 O

SR CA

LC STN Files: CA, CAPLUS

$$\begin{array}{c} & & & \\ & &$$

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:294739

ANSWER 2 OF 314 REGISTRY COPYRIGHT 2002 ACS L17

408365-70-8 REGISTRY . RN

Carbamic acid, [2-(2-pyridinyl)ethyl][4-[[2-[[3-CN (trifluoromethyl)phenyl]amino]benzoyl]amino]phenyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

OTHER NAMES:

tert-Butyl 2-(2-pyridinyl)ethyl[4-[[2-[3-(trifluoromethyl)anilino]benzoyl]amino]phenyl]carbamate

FS 3D CONCORD

C32 H31 F3 N4 O3 MF

SR CA

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:294739

ANSWER 5 OF 314 REGISTRY COPYRIGHT 2002 ACS L17 408364-90-9 REGISTRY RN

> 308-4994 Shears Searcher :

CN Benzamide, N-[4-[[2-(2-pyridinyl)ethyl]amino]phenyl]-2-[[3-(trifluoromethyl)phenyl]amino]- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN N-[4-[[2-(2-Pyridinyl)ethyl]amino]phenyl]-2-[3-(trifluoromethyl)anilino]benzamide

FS 3D CONCORD

MF C27 H23 F3 N4 O

SR CA

t,

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:294739

L17 ANSWER 6 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 401905-99-5 REGISTRY

CN L-Phenylalanine, N-(2,6-dichlorobenzoyl)-4-[[2-(phenylamino)benzoyl]amino]-(9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C29 H23 C12 N3 O4

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:217047

L17 ANSWER 7 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 381240-33-1 REGISTRY

CN Cyclohexanecarboxamide, N-(cyanophenylmethyl)-2-(phenylamino)-,

(1R,2S)-rel- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C21 H23 N3 O

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

Relative stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:53544

L17 ANSWER 14 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 372118-10-0 REGISTRY

CN 2-Pyridinecarboxamide, 5-bromo-3-[[(3-chloro-4-methoxyphenyl)methyl]amino]-N-(2-pyrimidinylmethyl)- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H17 Br Cl N5 O2

SR CA

LC STN Files: CA, CAPLUS

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:357948

L17 ANSWER 15 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **372117-99-2** REGISTRY

CN 3-Pyridinecarboxamide, 6-chloro-2-[[(3-chloro-4-methoxyphenyl)methyl]amino]-N-(2-pyrimidinylmethyl)- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H17 C12 N5 O2

SR CA

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:357948

L17 ANSWER 16 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 372115-94-1 REGISTRY

CN 2-Pyridinecarboxamide, 3-[[(3-chloro-4-methoxyphenyl)methyl]amino]-5[(2S)-2-(hydroxymethyl)-1-pyrrolidinyl]-N-[2-(4-morpholinyl)ethyl](9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C25 H34 C1 N5 O4

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:357948

L17 ANSWER 23 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 358659-88-8 REGISTRY

CN Benzamide, N-(5-chloro-2-pyridinyl)-5-fluoro-2-[[[4-(imino-1-pyrrolidinylmethyl)phenyl]methyl]amino]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C24 H23 C1 F N5 O

SR CA

LC STN Files: CA, CAPLUS, USPATZ, USPATFULL

3 REFERENCES IN FILE CA (1967 TO DATE)

3 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:69743

REFERENCE 2: 135:226888

REFERENCE 3: 135:210946

L17 ANSWER 51 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **352228-00-3** REGISTRY

CN 3-Pyridinecarboxamide, 2-[[(4-hydroxyphenyl)methyl]amino]-N-[3-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C20 H16 F3 N3 O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:137407

L17 ANSWER 52 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **352227-92-0** REGISTRY

CN 3-Pyridinecarboxamide, 2-[[(3-hydroxyphenyl)methyl]amino]-N-[3-(trifluoromethyl)phenyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C20 H16 F3 N3 O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:137407

L17 ANSWER 54 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **351380-07-9** REGISTRY.

CN 3-Pyridinecarboxamide, 4-(1-ethylpropoxy)-N,N,6-trimethyl-2-[(2,4,6-trimethylphenyl)amino]- (9CI) (CA INDEX NAME)

MF C23 H33 N3 O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:137515

L17 ANSWER 55 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **340719-29-1** REGISTRY

CN L-Phenylalanine, 4-[[2-[(diethylamino)carbonyl]-6-nitrophenyl]amino]-N-[[1-[(dimethylamino)carbonyl]cyclopropyl]carbonyl]- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C27 H33 N5 O7

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:367194

L17 ANSWER 57 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 337360-72-2 REGISTRY

CN Benzamide, N-[(3-chloro-4-methoxyphenyl)methyl]-5-cyano-2-[[(1R,4S)-4-hydroxy-2-cyclopenten-1-yl]amino]- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C21 H20 C1 N3 O3

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:340354

L17 ANSWER 58 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **330785-12-1** REGISTRY

CN Pyrazinecarboxamide, 3-[[(3-chloro-4-methoxyphenyl)methyl]amino]-5-(5,6-dihydroimidazo[1,2-a]pyrazin-7(8H)-yl)-N-[(4-methyl-2-morpholinyl)methyl]-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C25 H31 C1 N8 O3

SR CA

LC STN Files: CA, CAPLUS

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:252363

L17 ANSWER 63 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 330784-45-7 REGISTRY

CN Pyrazinecarboxamide, 3-[[(3-chloro-4-methoxyphenyl)methyl]amino]-5[(2S)-2-(hydroxymethyl)-1-pyrrolidinyl]-N-[2-(4-morpholinyl)ethyl](9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C24 H33 C1 N6 O4

SR CA

LC STN Files: CA, CAPLUS

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:252363

L17 ANSWER 66 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 328124-74-9 REGISTRY

CN Carbamic acid, [[2,3-dihydro-3-[2-[[2-(phenylamino)benzoyl]amino]eth yl]-5-benzofuranyl]iminomethyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C29 H32 N4 O4

SR CA

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:207826

L17 ANSWER 67 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 328123-93-9 REGISTRY

CN Benzamide, N-[2-[5-(aminoiminomethyl)-2,3-dihydro-3-benzofuranyl]ethyl]-2-(phenylamino)- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C24 H24 N4 O2

SR CA

LC STN Files: CA, CAPLUS

$$H_2N-C$$
 $H_2N-CH_2-CH_2-NH-C$
 $H_2N-CH_2-CH_2-NH-C$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:207826

L17 ANSWER 68 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **321438-66-8** REGISTRY

CN Benzamide, 2-[(2-ethyl-4-iodophenyl)amino]-N-(2-hydroxyethyl)-5-nitro-(9CI) (CA INDEX NAME)

OTHER NAMES:

CN N-(2-Hydroxyethyl)-2-(4-iodo-2-ethylphenylamino)-5-nitrobenzamide

FS 3D CONCORD

MF C17 H18 I N3 O4

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

$$\begin{array}{c|c} \mathsf{O} & & \mathsf{O} \\ \mathsf{HO-CH_2-CH_2-NH-C} & & \mathsf{Et} \\ & & \mathsf{NH-C} & & \mathsf{NH-C} \\ & & \mathsf{O_2N} & & \mathsf{I} \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:131317

L17 ANSWER 69 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 312324-33-7 REGISTRY

CN 1-Cyclohexene-1-carboxamide, 4,4-dimethyl-6-oxo-2[(phenylmethyl)amino]-N-[3-(trifluoromethyl)phenyl]- (9CI) (CA
INDEX NAME)

FS 3D CONCORD

MF C23 H23 F3 N2 O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

$$\begin{array}{c|c}
 & O & O \\
 & NH-C & O \\
 & Ph-CH_2-NH & Me
\end{array}$$
CF3

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:25114

L17 ANSWER 70 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 308832-00-0 REGISTRY

CN Carbamic acid, [(2,6-difluorophenyl)methyl][5-[4[[(methoxyamino)carbonyl]amino]phenyl]-3-[[[4(methoxymethoxy)phenyl]amino]carbonyl]-4[[methyl(phenylmethyl)amino]methyl]-2-thienyl]-, ethyl ester (9CI)
(CA INDEX NAME)

MF C40 H41 F2 N5 O7 S

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:4946

L17 ANSWER 71 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 308361-85-5 REGISTRY

CN Benzamide, 5-bromo-2-[[(3-hydroxy-4-methoxyphenyl)methyl]amino]-N-[2-[[[1-[(3-hydroxy-4-methoxyphenyl)methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C31 H37 Br N4 O6

SR CA

LC STN Files: CA, CAPLUS

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:173028

REFERENCE 2: 134:5154

L17 ANSWER 73 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 302958-78-7 REGISTRY

CN 5-Thiazolecarboxamide, 4-methyl-2-[[2-[[3-(trifluoromethyl)phenyl]amino]benzoyl]amino]-N-(2,4,6-trimethylphenyl)- (9CI) (CA INDEX NAME)

MF C28 H25 F3 N4 O2 S

SR CA

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 133:321878

L17 ANSWER 74 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 277335-40-7 REGISTRY

CN Benzamide, 5-bromo-2-[(2-ethyl-4-iodophenyl)amino]-N-[2-(1-pyrrolidinyl)ethyl]- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 5-Bromo-2-(4-iodo-2-ethylphenylamino)-N-(2-pyrrolidin-1-ylethyl)benzamide

FS 3D CONCORD

MF C21 H25 Br I N3 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:131317

REFERENCE 2: 133:73860

L17 ANSWER 75 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 277315-10-3 REGISTRY

CN Benzamide, 5-fluoro-2-[(4-iodo-2-methylphenyl)amino]-N-[2-(1-piperazinyl)ethyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C20 H24 F I N4 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

F
$$C-NH-CH_2-CH_2-N$$
NH
NH
NH

5 REFERENCES IN FILE CA (1967 TO DATE) 5 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 133:89333

REFERENCE 2: 133:89332

REFERENCE 3: 133:73860

REFERENCE 4: 133:73859

REFERENCE 5: 133:58616

L17 ANSWER 76 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **267891-63-4** REGISTRY

CN Benzamide, N-[2-(4-chlorophenyl)ethyl]-2-[[(4-hydroxyphenyl)methyl]amino]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C22 H21 C1 N2 O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 1 REFERENCES IN FILE CA (1967 TO DATE)
- 1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 132:334364

L17 ANSWER 79 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **267656-22-4** REGISTRY

CN 3-Pyridinecarboxamide, N-[5-(1-methylethyl)-2-thiazolyl]-2-[[3-(trifluoromethyl)phenyl]amino]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H17 F3 N4 O S

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 132:334454

L17 ANSWER 81 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 267405-06-1 REGISTRY

CN Benzamide, N-[[trans-4-[2-(methylamino)-2-oxoethyl]-1-

phenylcyclohexyl]methyl]-2-(phenylamino)- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C29 H33 N3 O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

Relative stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 132:334285

L17 ANSWER 82 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 262447-34-7 REGISTRY

CN Benzamide, 2-[(2,3-dimethylphenyl)amino]-N-[2-[4-[(imino-2-thienylmethyl)amino]phenyl]ethyl]- (9CI) (CA INDEX NAME)

MF C28 H28 N4 O S

CI COM

SR CA

LC STN Files: CA, CAPLUS

$$\begin{array}{c|c} \text{NH} & \text{O} & \text{O} \\ \text{C-NH} & \text{CH}_2\text{-}\text{CH}_2\text{-}\text{NH} - \text{C} \\ \text{NH} & \text{Me} \\ \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 132:251068

L17 ANSWER 84 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **261766-43-2** REGISTRY

CN Glycine, N-[2-[(2,6-dichloro-3-methylphenyl)amino]benzoyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C16 H14 C12 N2 O3

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:327362

REFERENCE 2: 132:231510

L17 ANSWER 91 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 254742-75-1 REGISTRY

CN 3-Pyridinecarboxamide, 2-[[[2'-[[(4,5-dimethyl-3-isoxazolyl)amino]sulfonyl][1,1'-biphenyl]-4-yl]methyl]propylamino]-N-methyl- (9CI) (CA INDEX NAME)

MF C28 H31 N5 O4 S

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:46172

REFERENCE 2: 132:93309

L17 ANSWER 92 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **254739-90-7** REGISTRY

CN 3-Pyridinecarboxamide, 2-[[[2'-[[(3,4-dimethyl-5-isoxazolyl)amino]sulfonyl][1,1'-biphenyl]-4-yl]methyl]propylamino]-N-methyl- (9CI) (CA INDEX NAME)

MF C28 H31 N5 O4 S

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER

2 REFERENCES IN FILE CA (1967 TO DATE)

2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 135:46172

REFERENCE 2: 132:93309

L17 ANSWER 93 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **252355-17-2** REGISTRY

CN Benzamide, 2,2'-iminobis[N-[3-(1-methyl-4-piperidinyl)-1H-indol-5-

yl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C42 H45 N7 O2

SR CA

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 1 REFERENCES IN FILE CA (1967 TO DATE)
- 1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 132:35606

L17 ANSWER 94 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 228581-72-4 REGISTRY

CN 1-Piperazinecarboxamide, 4-methyl-N-[[2-[[2-(phenylamino)-4-(phenylethynyl)benzoyl]amino]phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

MF C33 H31 N5 O4 S

SR CA

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 131:73440

L17 ANSWER 130 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 228580-98-1 REGISTRY

CN Benzoic acid, 2-[[2-(phenylamino)-5-(phenylethynyl)benzoyl]amino](9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C28 H20 N2 O3

SR CA

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 131:73440

L17 ANSWER 150 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 226245-19-8 REGISTRY

CN Benzamide, 5-bromo-2-[[(4-ethylphenyl)methyl]amino]-N-[2-[[(3R)-1-[(4-ethylphenyl)methyl]-3-pyrrolidinyl]amino]-2-oxoethyl]- (9CI) (CA INDEX NAME)

FS STEREOSEARCH

MF C31 H37 Br N4 O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3 REFERENCES IN FILE CA (1967 TO DATE)

3 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:263090

REFERENCE 2: 134:5154

REFERENCE 3: 131:18925

L17 ANSWER 151 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 226243-29-4 REGISTRY

CN Benzamide, 5-chloro-N-[2-oxo-2-[[[1-[(4-propylphenyl)methyl]-4-

piperidinyl]methyl]amino]ethyl]-2-[[(4-propylphenyl)methyl]amino](9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C35 H45 C1 N4 O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3 REFERENCES IN FILE CA (1967 TO DATE)

3 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:263090

REFERENCE 2: 134:5154

REFERENCE 3: 131:18925

L17 ANSWER 156 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 226241-83-4 REGISTRY

CN Benzamide, 5-bromo-2-[[(4-hydroxy-3-methoxyphenyl)methyl]amino]-N-[2-[[[1-[(4-hydroxy-3-methoxyphenyl)methyl]-4-piperidinyl]methyl]amino]-2-oxoethyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C31 H37 Br N4 O6

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3 REFERENCES IN FILE CA (1967 TO DATE)
3 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:263090

REFERENCE 2: 134:5154

REFERENCE 3: 131:18925

L17 ANSWER 155 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **226242-54-2** REGISTRY

CN Benzamide, 5-chloro-2-[[[4-(methylthio)phenyl]methyl]amino]-N-[2-[[[1-[[4-(methylthio)phenyl]methyl]-4-piperidinyl]methyl]amino]-2oxoethyl]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C31 H37 C1 N4 O2 S2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

MeS
$$CH_2 - NH - C - CH_2 - N$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3 REFERENCES IN FILE CA (1967 TO DATE)

3 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 136:263090

REFERENCE 2: 134:5154

REFERENCE 3: 131:18925

L17 ANSWER 167 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 223535-91-9 REGISTRY

CN [4,4'-Bipiperidine]-1-carboxylic acid, 1'-[4-[[(3-ethoxy-3-oxopropyl)amino]carbonyl]-3-[(phenylmethyl)amino]phenyl]-,
1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C34 H48 N4 O5

SR CA

LC STN Files: CA, CAPLUS

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 130:311817

L17 ANSWER 169 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **212630-37-0** REGISTRY

CN Benzamide, N-cyclohexyl-2-[(4-iodo-2-methylphenyl)amino]-5-nitro-

(9CI) (CA INDEX NAME)

OTHER NAMES:

CN N-Cyclohexyl-2-(4-iodo-2-methylphenylamino)-5-nitrobenzamide

FS 3D CONCORD

MF C20 H22 I N3 O3

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

8 REFERENCES IN FILE CA (1967 TO DATE)

8 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:131317

REFERENCE 2: 133:89333

REFERENCE 3: 133:89332

REFERENCE 4: 133:73860

REFERENCE 5: 133:73859

REFERENCE 6: 133:58616

REFERENCE 7: 130:124898

REFERENCE 8: 129:230537

L17 ANSWER 201 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 212629-93-1 REGISTRY

CN Benzamide, N-[2-[bis(1-methylethyl)amino]ethyl]-5-fluoro-2-[(4-iodo-2-methylphenyl)amino]- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN N-(2-Diisopropylaminoethyl)-5-fluoro-2-(4-iodo-2-methylphenylamino)benzamide

FS 3D CONCORD

MF C22 H29 F I N3 O

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

8 REFERENCES IN FILE CA (1967 TO DATE)

8 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:131317

REFERENCE 2: 133:89333

REFERENCE 3: 133:89332

REFERENCE 4: 133:73860

REFERENCE 5: 133:73859

REFERENCE 6: 133:58616

REFERENCE 7: 130:124898

REFERENCE 8: 129:230537

L17 ANSWER 280 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **212628-99-4** REGISTRY

CN Benzamide, 5-bromo-3,4-difluoro-N-(2-hydroxyethyl)-2-[(4-iodo-2-methylphenyl)amino]- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 5-Bromo-3, 4-difluoro-N-(2-hydroxyethyl)-2-(4-iodo-2-

methylphenylamino) benzamide

FS 3D CONCORD

MF C16 H14 Br F2 I N2 O2

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

$$\begin{array}{c|c} \mathsf{NH} & \mathsf{NH} \\ \mathsf{Br} & \mathsf{F} \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

8 REFERENCES IN FILE CA (1967 TO DATE)

8 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:131317

REFERENCE 2: 133:89333

REFERENCE 3: 133:89332

REFERENCE 4: 133:73860

REFERENCE 5: 133:73859

REFERENCE 6: 133:58616

REFERENCE 7: 130:124898

REFERENCE 8: 129:230537

L17 ANSWER 296 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 202827-87-0 REGISTRY

CN Benzamide, N-[4-(acetylamino)phenyl]-2-[(2,4,6-trinitrophenyl)amino]-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C21 H16 N6 O8

SR CAS Registry Services

LC STN Files: CA, CAPLUS, CHEMCATS, TOXCENTER

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 134:25114

L17 ANSWER 297 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 182434-95-3 REGISTRY

CN 9H-Fluorene-9-carboxamide, 9-[4-[4-[[2-(methylphenylamino)benzoyl]amino]-1-piperidinyl]butyl]-N-(2,2,2-trifluoroethyl)- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C39 H41 F3 N4 O2

CI COM

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PAGE 1-A

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 128:154011

REFERENCE 2: 125:275663

L17 ANSWER 298 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 182432-11-7 REGISTRY

CN 9H-Fluorene-9-carboxamide, N-(2,2,2-trifluoroethyl)-9-[4-[4-[[2-[[3-(trifluoromethyl)phenyl]amino]benzoyl]amino]-1-piperidinyl]butyl]-(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C39 H38 F6 N4 O2

CI COM

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PAGE 1-A

$$_{\mathrm{F_{3}C-CH_{2}-NH-C}}^{\mathrm{O}}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 128:154011

REFERENCE 2: 125:275663

L17 ANSWER 299 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 182431-88-5 REGISTRY

CN 3-Pyridinecarboxamide, N-[1-[4-[9-[[(2,2,2-trifluoroethyl)amino]carbonyl]-9H-fluoren-9-yl]butyl]-4-piperidinyl]-2-[[3-(trifluoromethyl)phenyl]amino]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C38 H37 F6 N5 O2

CI COM

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PAGE 1-A

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 128:154011

REFERENCE 2: 125:275663

L17 ANSWER 300 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 182429-79-4 REGISTRY

CN 9H-Fluorene-9-carboxamide, 9-[4-[4-[[2-[(phenylmethyl)amino]benzoyl] amino]-1-piperidinyl]butyl]-N-(2,2,2-trifluoroethyl)- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C39 H41 F3 N4 O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PAGE 1-A

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 128:282780

REFERENCE 2: 125:275663

L17 ANSWER 301 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 182429-76-1 REGISTRY

CN 9H-Fluorene-9-carboxamide, 9-[4-[4-[[2-(phenylamino)benzoyl]amino]-1-piperidinyl]butyl]-N-(2,2,2-trifluoroethyl)- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C38 H39 F3 N4 O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PAGE 1-A

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 128:282780

REFERENCE 2: 125:275663

L17 ANSWER 302 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 169317-90-2 REGISTRY

CN Propanedioic acid, methoxy[[[1-(phenylmethyl)-5[(phenylmethyl)amino]-1H-imidazol-4-yl]carbonyl]amino]-, diethyl
ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C26 H30 N4 O6

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)
2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 130:38644

REFERENCE 2: 123:286534

L17 ANSWER 303 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **169043-59-8** REGISTRY

CN Benzoic acid, 4-[[[2-[[(1,3-benzodioxol-5-ylmethyl)amino]carbonyl]-4-chlorophenyl]amino]methyl]-, methyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C24 H21 C1 N2 O5

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 123:256357

L17 ANSWER 304 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 163267-27-4 REGISTRY

CN Benzamide, N-[1-(3,3-diphenylpropyl)-4-piperidinyl]-2-(phenylamino)-

(9CI) (CA INDEX NAME)
FS 3D CONCORD

MF C33 H35 N3 O

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)

2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 128:282780

REFERENCE 2: 123:169516

L17 ANSWER 305 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **159619-35-9** REGISTRY

CN Benzamide, 5-chloro-N-[2-(diethylamino)ethyl]-2[methyl(phenylmethyl)amino]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C21 H28 C1 N3 O

CI COM

SR CA

LC STN Files: CA, CAPLUS

$$\begin{array}{c|c} & \text{C1} \\ \hline \\ \text{C-NH-CH}_2\text{-CH}_2\text{-NEt}_2 \\ \hline \\ \text{Ph-CH}_2\text{-N} \\ \text{O} \\ \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)

2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 125:265033

REFERENCE 2: 122:31131

L17 ANSWER 306 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 152814-88-5 REGISTRY

FS 3D CONCORD

MF C21 H19 N3 O2

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 120:164213

L17 ANSWER 307 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 150780-12-4 REGISTRY

CN Cyclohexanecarbothioamide, 1-imidazo[1,2-a]pyridin-6-yl-N-methyl-2-[(phenylmethyl)amino]- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Imidazo[1,2-a]pyridine, cyclohexanecarbothioamide deriv.

MF C22 H26 N4 S

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 119:225951

L17 ANSWER 308 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN 136148-82-8 REGISTRY

CN Glycine, N-[2-[[(2,4-dichlorophenyl)methyl]amino]-5-methoxybenzoyl]-, ethyl ester (9CI) (CA INDEX NAME)

FS 3D CONCORD

DR 136148-79-3

MF C19 H20 C12 N2 O4

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 115:159174

L17 ANSWER 309 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **133670-75-4** REGISTRY

CN Cyclohexanecarbothioamide, N-methyl-2-(phenylamino)-1-(3-pyridinyl), trans- (9CI) (CA INDEX NAME)

, trans- (9CI) (CA OTHER CA INDEX NAMES:

CN Cyclohexanecarbothioamide, N-methyl-2-(phenylamino)-1-(3-pyridinyl)-, trans-(.+-.)-

STEREOSEARCH FS

C19 H23 N3 S MF

SR CA

LC STN Files: CA, CAPLUS, USPATFULL

Relative stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 114:207038

L17 ANSWER 310 OF 314 REGISTRY COPYRIGHT 2002 ACS

133667-59-1 REGISTRY RN

Cyclohexanecarbothioamide, N-methyl-2-[(phenylmethyl)amino]-1-(3-CN

pyridinyl)-, trans- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

Cyclohexanecarbothioamide, N-methyl-2-[(phenylmethyl)amino]-1-(3-CN

pyridinyl)-, trans-(.+-.)-

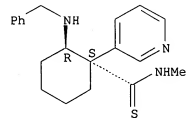
FS STEREOSEARCH

C20 H25 N3 S MF

SR CA

CA, CAPLUS, USPATFULL LC STN Files:

Relative stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 114:207038

> Shears 308-4994 Searcher :

L17 ANSWER 312 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **89459-32-5** REGISTRY

CN Benzamide, 2-[(2-acetylphenyl)amino]-N-[2-(dimethylamino)ethyl]-

(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C19 H23 N3 O2

LC STN Files: BEILSTEIN*, CA, CAPLUS, CASREACT, TOXCENTER, USPATFULL

(*File contains numerically searchable property data)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

2 REFERENCES IN FILE CA (1967 TO DATE)

2 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 106:156252

REFERENCE 2: 100:138978

L17 ANSWER 313 OF 314 REGISTRY COPYRIGHT 2002 ACS

RN **71908-19-5** REGISTRY

CN Benzamide, N-[2-(nitrooxy)ethyl]-2-[[3-(trifluoromethyl)phenyl]amino

]- (9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C16 H14 F3 N3 O4

LC STN Files: CA, CAPLUS

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 91:192973
FILE °CAOLD° ENTERED AT 14:53:51 ON 31 MAY 2002

L18 0 S L17 "OSPATEULL' ENTERED AT 14:53:58 ON 31 MAY 2002) L19 26 SEA ABB=ON PLU=ON L17 26 SEA ABB=ON PLU=ON L19 AND (PROPHYLACT? OR PROPHYLAX? L20 OR TREAT? OR THERAP?) 25 SEA ABB=ON PLU=ON L20 AND (DISEAS? OR DISORDER OR L21 MALAD?) L21 ANSWER 1 OF 25 USPATFULL ACCESSION NUMBER: 2002:63897 USPATFULL TITLE: Cyclic amine derivatives and their use as drugs Shiota, Tatsuki, Hino, JAPAN INVENTOR(S): Kataoka, Ken-ichiro, Hino, JAPAN Imai, Minoru, Hino, JAPAN Tsutsumi, Takaharu, Hino, JAPAN Sudoh, Masaki, Handa, JAPAN Sogawa, Ryo, Hino, JAPAN Morita, Takuya, Hino, JAPAN Hada, Takahiko, Okayama, JAPAN Muroga, Yumiko, Hino, JAPAN Takenouchi, Osami, Hino, JAPAN Furuya, Minoru, Hino, JAPAN Endo, Noriaki, Hino, JAPAN Tarby, Christine M., Wilmington, DE, United States Moree, Wilna, San Diego, CA, United States Teig, Steven, Palo Alto, CA, United States PATENT ASSIGNEE(S): Teijin Limited, Osaka, JAPAN (non-U.S. corporation) Dupont Pharmaceuticals Research Laboratories, San Diego, CA, United States (U.S. corporation) NUMBER KIND DATE PATENT INFORMATION: US 6362177 В1 20020326 US 2001-905078 20010716 APPLICATION INFO.: Division of Ser. No. US 2000-554562, filed on 16 RELATED APPLN. INFO.: May 2000 DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED PRIMARY EXAMINER: Aulakh, Charanjit S. LEGAL REPRESENTATIVE: Sughrue Mion, PLLC NUMBER OF CLAIMS: 22 EXEMPLARY CLAIM: NUMBER OF DRAWINGS: 0 Drawing Figure(s); 0 Drawing Page(s) LINE COUNT: 7859 CAS INDEXING IS AVAILABLE FOR THIS PATENT. AB ##STR1## A compound represented by the general formula (I), a pharmaceutically acceptable acid addition salt thereof or a

A compound represented by the general formula (I), a pharmaceutically acceptable acid addition salt thereof or a pharmaceutically acceptable C.sub.1-C.sub.6 alkyl addition salt thereof, and their medical applications. These compounds inhibit the action of chemokines such as MIP-1.alpha. and/or MCP-1 on target cells, and are useful as therapeutic and/or preventative drugs in diseases, such as atheroclerosis, rheumatoid arthritis, and the like where blood monocytes and lymphocytes infiltrate into tissues.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 2 OF 25 USPATFULL

ACCESSION NUMBER:

2002:37927 USPATFULL

TITLE:

2-(4-bromo or 4-iodo phenylamino) benzoic acid

derivatives

INVENTOR(S):

Barrett, Stephen Douglas, Livonia, MI, UNITED

STATES

Bridges, Alexander James, Saline, MI, UNITED

STATES

Cody, Donna Reynolds, Saline, MI, UNITED STATES

Doherty, Annette Marian, Paris, FRANCE Dudley, David Thomas, Ann Arbor, MI, UNITED

STATES

Saltiel, Alan Robert, Ann Arbor, MI, UNITED

STATES

Schroeder, Mel Conrad, Dexter, MI, UNITED STATES

Tecle, Haile, Ann Arbor, MI, UNITED STATES

KIND DATE NUMBER _____ 20020221 A1

PATENT INFORMATION: APPLICATION INFO.:

US 2002022647

RELATED APPLN. INFO.:

US 2001-931596 A1 20010816 (9)

Division of Ser. No. US 2000-462319, filed on 5 Jan 2000, GRANTED, Pat. No. US 6310060 A 371 of International Ser. No. WO 1998-US13105, filed on

24 Jun 1998, UNKNOWN

NUMBER DATE ______

PRIORITY INFORMATION:

US 1997-60051433 19970701 19970701 (60)

US 1997~51433P Utility

DOCUMENT TYPE: FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE:

Warner-Lambert Company, 2800 Plymouth Road, Ann

Arbor, MI, 48105

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

34 1

LINE COUNT:

1564

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Phenylamino benzoic acid, benzamides, and benzyl alcohol AB

> derivatives of the formula ##STR1##

where R.sub.1, R.sub.2, R.sub.3, R.sub.4, R.sub.5, and R.sub.6 are hydrogen or substituent groups such as alkyl, and where R.sub.7 is hydrogen or an organic radical, and Z is COOR.sub.7,

tetrazolyl, CONR.sub.6R.sub.7, or CH.sub.20R.sub.7, are potent inhibitors of MEK and, as such, are effective in treating cancer and other proliferative diseases such as inflammation, psoriasis and restenosis, as well as stroke, heart failure, and immunodeficiency disorders.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 3 OF 25 USPATFULL

ACCESSION NUMBER:

2002:32592 USPATFULL

TITLE:

N-aryl(thio)anthranilic acid amide derivatives,

Searcher : 308-4994 Shears

their preparation and their use as VEGF receptor

tyrosine kinase inhibitors

Altmann, Karl-Heinz, Reinach, SWITZERLAND INVENTOR(S): Bold, Guido, Gipf-Oberfrick, SWITZERLAND

Furet, Pascal, Thann, FRANCE

Manley, Paul William, Arlesheim, SWITZERLAND Wood, Jeanette Marjorie, Biel-Benken, SWITZERLAND

Ferrari, Stefano, Muttenz, SWITZERLAND Hofmann, Francesco, Bottmingen, SWITZERLAND Mestan, Jurgen, Denzlingen, GERMANY, FEDERAL

REPUBLIC OF

Huth, Andreas, Berlin, GERMANY, FEDERAL REPUBLIC

Kruger, Martin, Berlin, GERMANY, FEDERAL REPUBLIC

Seidelmann, Dieter, Berlin, GERMANY, FEDERAL

REPUBLIC OF

Menrad, Andreas, Oranienburg, GERMANY, FEDERAL

REPUBLIC OF

Haberey, Martin, Berlin, GERMANY, FEDERAL

REPUBLIC OF

Thierauch, Karl-Heinz, Berlin, GERMANY, FEDERAL

REPUBLIC OF

NUMBER KIND DATE _______

PATENT INFORMATION: APPLICATION INFO.:

US 2002019414 A1 20020214 US 2001-850434 A1 20010507 (9)

RELATED APPLN. INFO.:

Continuation of Ser. No. WO 1999-EP8545, filed on

8 Nov 1999, UNKNOWN

NUMBER DATE _____

PRIORITY INFORMATION:

GB 1998-24579 19981110

DOCUMENT TYPE: FILE SEGMENT:

Utility APPLICATION

LEGAL REPRESENTATIVE:

THOMAS HOXIE, NOVARTIS CORPORATION, PATENT AND

TRADEMARK DEPT, 564 MORRIS AVENUE, SUMMIT, NJ,

079011027

NUMBER OF CLAIMS:

17 EXEMPLARY CLAIM:

LINE COUNT: 2620 . CAS INDEXING IS AVAILABLE FOR THIS PATENT.

##STR1## AB

> Described are compounds of formula (I), wherein W is O or S; X is NR.sub.8; Y is CR.sub.9R.sub.10-(CH.sub.2)n wherein R.sub.9 and R.sub.10 are independently of each other hydrogen or lower alkyl, and n is an integer of from and including 0 to and including 3; or Y is SO.sub.2; R.sub.1 is aryl; R.sub.2 is a mono- or bicyclic heteroaryl group comprising one or more ring nitrogen atoms with the exception that R.sub.2 cannot represent 2-phthalimidyl, and in case of Y.dbd.SO.sub.2 cannot represent 2,1,3-benzothiadiazol-4yl; any of R.sub.3, R.sub.4, R.sub.5 and R.sub.6, independently of the other, is H or a substituent other than hydrogen; and R.sub.7 and R.sub.8, independently of each other, are H or lower alkyl; or a N-oxide or a pharmaceutically acceptable salt thereof for the preparation of a pharmaceutical product for the treatment

of a neoplastic **disease** which responds to an inhibition of the VEGF receptor tyrosine kinase activity. The compounds of formula (I) can be used for the **treatment** e.g. of a neoplastic **disease**, such as a tumor **disease**, of retinopathy and age-related macular degeneration.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 4 OF 25 USPATFULL

ACCESSION NUMBER: 2002:27513 USPATFULL

TITLE: Beta-amino acid nitrile derivatives as cathepsin

K inhibitors

INVENTOR(S): Gabriel, Tobias, Loerrach, GERMANY, FEDERAL

REPUBLIC OF

Pech, Michael, Hartheim, GERMANY, FEDERAL

REPUBLIC OF

Rodriguez Sarmiento, Rosa Maria, Basle,

SWITZERLAND

NUMBER DATE

PRIORITY INFORMATION: EP 2000-112577 20000614

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HOFFMANN-LA ROCHE INC., PATENT LAW DEPARTMENT,

340 KINGSLAND STREET, NUTLEY, NJ, 07110

NUMBER OF CLAIMS: 294
EXEMPLARY CLAIM: 1
LINE COUNT: 3380

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to beta-amino acid nitrile derivatives and pharmaceutically acceptable salts and/or pharmaceutically acceptable esters thereof. The compounds are cysteine protease inhibitors useful for the treatment of diseases associated with cysteine proteases, such as osteoporosis, osteoarthritis, rheumatoid arthritis, tumor metastasis, glomerulonephritis, atherosclerosis, myocardial infarction, angina pectoris, instable angina pectoris, stroke, plaque rupture, transient ischemic attacks, amaurosis fugax, peripheral arterial occlusive disease, restenosis after angioplasty and stent placement, abdominal aortic aneurysm formation, inflammation, autoimmune disease, malaria, ocular fundus tissue cytopathy and respiratory disease.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 5 OF 25 USPATFULL

ACCESSION NUMBER: 2002:27480 USPATFULL

TITLE: Corticotropin releasing factor antagonists
INVENTOR(S): Chen, Yuhpyng L., Waterford, CT, UNITED STATES

NUMBER KIND DATE

PATENT INFORMATION: US 2002016328 A1 20020207 APPLICATION INFO.: US 2001-761995 A1 20010117 (9)

NUMBER DATE

PRIORITY INFORMATION: US 2000-176611P 20000118 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: PFIZER INC, 150 EAST 42ND STREET, 5TH FLOOR -

STOP 49, NEW YORK, NY, 10017-5612

NUMBER OF CLAIMS: 22 EXEMPLARY CLAIM: 1 LINE COUNT: 5425

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Corticotropin-releasing factor (CRF) antagonists having the

formulae ##STR1##

wherein the dashed lines, A, B, Y, Z, G, R.sub.3, R.sub.4, R.sub.5, R.sub.6, R.sub.16 and R.sub.17 are as defined in the application, and processes for preparing them. These compounds and their pharmaceutically acceptable salts are useful in the treatment disorders including CNS and

stress-related disorders.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 6 OF 25 USPATFULL

ACCESSION NUMBER: 2002:14003 USPATFULL

TITLE: Thienopyrimidine compounds, their production and

use

INVENTOR(S): Furuya, Shuichi, Tsukuba, JAPAN

Suzuki, Nobuhiro, Tsukuba, JAPAN Choh, Nobuo, Tsukuba, JAPAN Nara, Yoshi, Suita, JAPAN

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Osaka, JAPAN

(non-U.S. corporation)

NUMBER DATE

PRIORITY INFORMATION: JP 1999-79371 19990324 JP 2000-18019 20000125

DOCUMENT TYPE: Utility
FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Ford, John M.

LEGAL REPRESENTATIVE: Chao, Mark, Ramesh, Elaine M.

NUMBER OF CLAIMS: 24
EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 1 Drawing Figure(s); 1 Drawing Page(s)

LINE COUNT: 1944

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A compound of the formula: ##STR1##

wherein R.sup.1 and R.sup.2 each is hydrogen, hydroxy, C.sub.1-4 alkoxy, C.sub.1-4 alkoxy-carbonyl or C.sub.1-4 alkyl which may be substituted; R.sup.3 is hydrogen, halogen, hydroxy or C.sub.1-4 alkoxy which may be substituted; or adjacent two R.sup.3 may form C.sub.1-4 alkylenedioxy; R.sup.4 is hydrogen or C.sub.1-4 alkyl; R.sup.6 is C.sub.1-4 alkyl which may be substituted or a group of the formula: ##STR2##

wherein R.sup.5 is hydrogen or R.sup.4 and R.sup.5 may form heterocycle; and n is 0-5, or a salt thereof, has an excellent GnRH-antagonizing activity, and is useful for preventing or treating sex hormone-dependent diseases.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 7 OF 25 USPATFULL

ACCESSION NUMBER: 2002:4204 USPATFULL

TITLE: Benzamides and related inhibitors of factor Xa INVENTOR(S): Zhu, Bing-Yan, Belmont, CA, UNITED STATES

Zhang, Penglie, Foster City, CA, UNITED STATES Wang, Lingyan, Chatham, NJ, UNITED STATES

Huang, Wenrong, Cupertino, CA, UNITED STATES Goldman, Erick A., San Francisco, CA, UNITED

STATES

Li, Wenhao, South San Francisco, CA, UNITED

STATES

Zuckett, Jingmei, Glendale, AZ, UNITED STATES Song, Yonghong, Foster City, CA, UNITED STATES Scarborough, Robert, Half Moon Bay, CA, UNITED

20000229 (60)

STATES

APPLICATION INFO.: US 2001-794225 A1 20010228 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 2000-663420,

filed on 15 Sep 2000, PENDING

NUMBER DATE

PRIORITY INFORMATION: US 2000-185746P
DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: MORGAN, LEWIS & BOCKIUS, 1800 M STREET NW,

WASHINGTON, DC, 20036-5869

NUMBER OF CLAIMS: 72
EXEMPLARY CLAIM: 1
LINE COUNT: 5918

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Novel benzamide compounds including their pharmaceutically acceptable isomers, salts, hydrates, solvates and prodrug derivatives having activity against mammalian factor Xa are described. Compositions containing such compounds are also described. The compounds and compositions are useful in vitro or in vivo for preventing or treating coagulation disorders.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 8 OF 25 USPATFULL

ACCESSION NUMBER:

2001:191130 USPATFULL

2-(4-bromo or 4-iodo phenylamino) benzoic acid TITLE: derivatives and their use as MEK inhibitors

Barrett, Stephen Douglas, Livonia, MI, United INVENTOR(S):

States

Bridges, Alexander James, Saline, MI, United

States

Cody, Donna Reynolds, Saline, MI, United States

Doherty, Annette Marian, Paris, France Dudley, David Thomas, Ann Arbor, MI, United

States

Saltiel, Alan Robert, Ann Arbor, MI, United

States

Schroeder, Mel Conrad, Dexter, MI, United States

Tecle, Haile, Ann Arbor, MI, United States

Warner-Lambert Company, Morris Plains, NJ, United PATENT ASSIGNEE(S):

States (U.S. corporation)

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 6310060 WO 9901421 US 2000-462319 WO 1998-US13105	B1	20011030 19990114 20000105 19980624 20000105	(9) PCT 371 date
			20000105	PCT 102(e) date

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

Higel, Floyd D. PRIMARY EXAMINER: Sackey, Ebenezer ASSISTANT EXAMINER: Ashbrook, Charles W. LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 37 EXEMPLARY CLAIM: 1 LINE COUNT: 1524

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Phenylamino benzoic acid, benzamides, and benzyl alcohol AΒ derivatives of the formula ##STR1##

> where R.sub.1, R.sub.2, R.sub.3, R.sub.4, R.sub.5, and R.sub.6 are hydrogen or substituent groups such as alkyl, and where R.sub.7 is hydrogen or an organic radical, and Z is COOR.sub.7, tetrazolyl, CONR.sub.6 R.sub.7, or CH.sub.2 OR.sub.7, are potent inhibitors of MEK and, as such, are effective in treating cancer and other proliferative diseases such as inflammation, psoriasis and restenosis, as well as stroke, heart failure, and immunodeficiency disorders.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 9 OF 25 USPATFULL

2001:168259 USPATFULL ACCESSION NUMBER:

Thienopyrimidine compounds, their production and TITLE:

Furuya, Shuichi, Ibaraki, Japan INVENTOR(S):

Suzuki, Nobuhiro, Ibaraki, Japan

308-4994 Searcher : Shears

Choh, Nobuo, Ibaraki, Japan

Nara, Yoshi, Osaka, Japan

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Osaka, Japan

(non-U.S. corporation)

NUMBER KIND DATE _____ US 6297379 B1 20011002 PATENT INFORMATION: 20000928 WO 2000056739 20000426 (9) US 2000-530495 APPLICATION INFO.: WO 2000-JP1777 20000323

20000426 PCT 371 date 20000426 PCT 102(e) date

DATE NUMBER ______

JP 1999-79371 19990324 PRIORITY INFORMATION:

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

PRIMARY EXAMINER: Shah, Mukund J. ASSISTANT EXAMINER: Rao, Deepak R.

LEGAL REPRESENTATIVE: Riesen, Philippe Y., Chao, Mark

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: . 1 Drawing Figure(s); 1 Drawing Page(s)

LINE COUNT: 1679

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB ##STR1##

> A compound of formula (I) wherein R.sup.1 and R.sup.2 each is hydrogen, hydroxy, C.sub.1-4 alkoxy, C.sub.1-4 alkoxy-carbonyl or C.sub.1-4 alkyl which may be substituted; R.sup.3 is hydrogen, halogen, hydroxy or C.sub.1-4 alkoxy which may be substituted; or adjacent two R.sup.3 may form C.sub.1-4 alkylenedioxy; R.sup.4 is hydrogen or C.sub.1-4 alkyl; R.sup.6 is C.sub.1-4 alkyl which may be substituted or a group of the formula (A) wherein R.sup.5 is hydrogen of R.sup.4 and R.sup.5 may form heterocycle; and n is 0-5, or a salt thereof, has an excellent GnRH-antagonizing activity, and is useful for preventing or treating sex hormone-dependent diseases.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 10 OF 25 USPATFULL

2000:64876 USPATFULL ACCESSION NUMBER:

Inhibitors of microsomal triglyceride transfer TITLE:

protein and method

Biller, Scott A., Hopewell, NJ, United States INVENTOR(S):

Dickson, John K., Eastampton, NJ, United States Lawrence, R. Michael, Yardley, PA, United States Magnin, David R., Hamilton, NJ, United States Poss, Michael A., Lawrenceville, NJ, United

States

Sulsky, Richard B., Franklin Park, NJ, United

States

Tino, Joseph A., Lawrenceville, NJ, United States Lawson, John E., Wallingford, CT, United States Holava, Henry M., Meriden, CT, United States

308-4994 Searcher : Shears

Partyka, Richard A., Neshanic, NJ, United States Bristol-Myers Squibb Company, Princeton, NJ, PATENT ASSIGNEE(S):

United States (U.S. corporation)

NUMBER KIND DATE _____ US 1997-898303 Continuet 20000523 PATENT INFORMATION: 19970721 (8) APPLICATION INFO.:

Continuation of Ser. No. US 1995-472067, filed on RELATED APPLN. INFO.:

6 Jun 1995, now patented, Pat. No. US 5739135 which is a continuation-in-part of Ser. No. US 1995-391901, filed on 21 Feb 1995, now abandoned which is a continuation-in-part of Ser. No. US 1994-284808, filed on 5 Aug 1994, now abandoned which is a continuation-in-part of Ser. No. US 1993-117362, filed on 3 Sep 1993, now patented,

Pat. No. US 5595872

DOCUMENT TYPE: Utility Granted FILE SEGMENT:

PRIMARY EXAMINER: Raymond, Richard L. ASSISTANT EXAMINER: Coleman, Brenda Rodney, Burton LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT: 5783

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compounds are provided which inhibit microsomal triglyceride transfer protein and thus are useful for lowering serum lipids and treating atherosclerosis and related diseases.

The compounds have the structure ##STR1## wherein R.sup.1 to R.sup.7, Q, X and Y are as defined herein.

CAS INDEXING IS AVAILABLE FOR THIS PATENT'.

L21 ANSWER 11 OF 25 USPATFULL

ACCESSION NUMBER: 2000:27994 USPATFULL

Inhibitors of microsomal triglyceride transfer TITLE:

protein and method

INVENTOR(S): Biller, Scott A., Hopewell, NJ, United States Dickson, John K., Eastampton, NJ, United States Lawrence, R. Michael, Yardley, PA, United States Magnin, David R., Hamilton, NJ, United States

Poss, Michael A., Lawrenceville, NJ, United

States

Sulsky, Richard B., Franklin Park, NJ, United

Tino, Joseph A., Lawrenceville, NJ, United States Lawson, John E., Wallingford, CT, United States Holava, Henry M., Meriden, CT, United States Partyka, Richard A., Neshanic, NJ, United States

Bristol-Myers Squibb Company, Princeton, NJ, PATENT ASSIGNEE(S):

United States (U.S. corporation)

NUMBER KIND DATE _______ US 1997-898304 Division 20000307 PATENT INFORMATION: 19970721 (8) APPLICATION INFO.: Division of Ser. No. US 1995-472067, filed on 6 RELATED APPLN. INFO.:

Jun 1995, now patented, Pat. No. US 5739135 which

is a continuation-in-part of Ser. No. US

1995-391901, filed on 21 Feb 1995, now abandoned which is a continuation-in-part of Ser. No. US 1994-284808, filed on 5 Aug 1994, now abandoned which is a continuation-in-part of Ser. No. US 1993-117362, filed on 3 Sep 1993, now patented,

Pat. No. US 5595872

DOCUMENT TYPE:

Utility Granted

FILE SEGMENT: PRIMARY EXAMINER:

Raymond, Richard L. Coleman, Brenda Rodney, Burton

LEGAL REPRESENTATIVE: NUMBER OF CLAIMS:

ASSISTANT EXAMINER:

EXEMPLARY CLAIM: LINE COUNT:

5940

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compounds are provided which inhibit microsomal triglyceride transfer protein and thus are useful for lowering serum lipids and

treating atherosclerosis and related diseases.

The compounds have the structure ##STR1## wherein R.sup.1 to

R.sup.7, Q, X and Y are as defined herein.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 12 OF 25 USPATFULL

ACCESSION NUMBER: 1999:34002 USPATFULL

TITLE:

Inhibitors of microsomal triglyceride transfer

protein and method

INVENTOR(S):

Biller, Scott A., Hopewell, NJ, United States Dickson, John K., Eastampton, NJ, United States Lawrence, R. Michael, Yardley, PA, United States Magnin, David R., Hamilton, NJ, United States Poss, Michael A., Lawrenceville, NJ, United

States

Sulsky, Richard B., Franklin Park, NJ, United

Tino, Joseph A., Lawrenceville, NJ, United States Lawson, John E., Wallingford, CT, United States Holava, Henry M., Meriden, CT, United States Partyka, Richard A., Neshanic, NJ, United States

PATENT ASSIGNEE(S):

Bristol-Myers Squibb Company, Princeton, NJ,

United States (U.S. corporation)

NUMBER KIND DATE ______ US 5883099 19990316

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

US 1997-896872 19970721 (8) Division of Ser. No. US 1995-472067, filed on 6

Jun 1995, now patented, Pat. No. US 5739135 which

is a continuation-in-part of Ser. No. US 1995-391901, filed on 21 Feb 1995, now abandoned which is a continuation-in-part of Ser. No. US 1994-284808, filed on 5 Aug 1994, now abandoned which is a continuation-in-part of Ser. No. US

1993-117362, filed on 3 Sep 1993, now patented, Pat. No. US 5595872

DOCUMENT TYPE:

Utility

308-4994 Shears Searcher :

FILE SEGMENT: Granted

PRIMARY EXAMINER: Huang, Evelyn LEGAL REPRESENTATIVE: Rodney, Burton

NUMBER OF CLAIMS: 9
EXEMPLARY CLAIM: 1
LINE COUNT: 5860

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Compounds are provided which inhibit microsomal triglyceride

transfer protein and thus are useful for lowering serum lipids and

treating atherosclerosis and related diseases.

The compounds have the structure ##STR1## wherein R.sup.1 to

R.sup.7, Q, X and Y are as defined herein.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 13 OF 25 USPATFULL

ACCESSION NUMBER: 1998:150915 USPATFULL

TITLE: Ring-expanded nucleosides and nucleotides

INVENTOR(S): Hosmane, Ramachandra, Columbia, MD, United States

Burns, Barry, Owings Mills, MD, United States

PATENT ASSIGNEE(S): Universy of Maryland, Baltimore, MD, United

States (U.S. corporation)

Nabi, Boca Raton, FL, United States (U.S.

corporation)

PATENT INFORMATION: US 5843912 19981201 APPLICATION INFO.: US 1995-518278 19950823 (8)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1994-268570,

filed on 6 Jul 1994, now abandoned

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Kunz, Gary L.

LEGAL REPRESENTATIVE: Cushman, Darby & Cushman, IP Group of Pillsbury,

Madison & Sutro LLP

NUMBER OF CLAIMS: 16 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 7 Drawing Figure(s); 5 Drawing Page(s)

LINE COUNT: 1891

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention relates to compositions comprising analogues of purine nucleosides containing a ring-expanded ("fat")

heterocyclic ring, in place of purine, and an unmodified or

modified sugar residue, pharmaceutically acceptable derivatives of

such compositions, as well as methods of use thereof. In particular, these compositions may be utilized in the

treatment of certain cancers, bacterial, fungal,

parasitic, and viral infections, including, but not limited to,

Acquired Immunodeficiency Syndrome (AIDS) and hepatitis.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 14 OF 25 USPATFULL

ACCESSION NUMBER: 1998:91828 USPATFULL

TITLE: Microsomal triglyceride transfer protein INVENTOR(S): Wetterau, II, John R., Langhorne, PA, United

States

Sharp, Daru Young, Perrineville, NJ, United

States

Gregg, Richard E., Pennington, NJ, United States E. R. Squibb & Sons, Inc., Princeton, NJ, United PATENT ASSIGNEE(S):

States (U.S. corporation)

NUMBER KIND DATE _____

US 5789197 19980804 PATENT INFORMATION: US 1995-486924 19950607 (8) APPLICATION INFO.:

Division of Ser. No. US 1993-117362, filed on 3 RELATED APPLN. INFO.:

Sep 1993, now patented, Pat. No. US 5595872 which

is a continuation-in-part of Ser. No. US

1993-15449, filed on 22 Feb 1993, now abandoned which is a continuation-in-part of Ser. No. US 1992-847503, filed on 6 Mar 1992, now abandoned

DOCUMENT TYPE: Utility Granted FILE SEGMENT:

Elliott, George C. PRIMARY EXAMINER: ASSISTANT EXAMINER: M'Garry, Sean

Gaul, Timothy J., Bogden, James M. LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

14 Drawing Figure(s); 8 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 4815

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Nucleic acid sequences, particularly DNA sequences, coding for all AB or part of the high molecular weight subunit of microsomal triglyceride transfer protein, expression vectors containing the DNA sequences, host cells containing the expression vectors, and methods utilizing these materials. The invention also concerns polypeptide molecules comprising all or part of the high molecular weight subunit of microsomal triglyceride transfer protein, and methods for producing these polypeptide molecules. The invention additionally concerns novel methods for preventing, stabilizing or causing regression of atherosclerosis and therapeutic agents having such activity. The invention concerns further novel methods for lowering serum liquid levels and therapeutic agents having such activity.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 15 OF 25 USPATFULL

1998:39526 USPATFULL ACCESSION NUMBER:

Inhibitors of microsomal triglyceride transfer TITLE:

protein and method

Biller, Scott A., Hopewell, NJ, United States INVENTOR(S):

Dickson, John K., Eastampton, NJ, United States Lawrence, R. Michael, Yardley, PA, United States Magnin, David R., Hamilton, NJ, United States Poss, Michael A., Lawrenceville, NJ, United

States

Sulsky, Richard B., Franklin Park, NJ, United

States

Tino, Joseph A., Lawrenceville, NJ, United States

Bristol-Myers Squibb Company, Princeton, NJ, PATENT ASSIGNEE(S):

United States (U.S. corporation)

Shears 308-4994 Searcher :

	NUMBER KIND DATE				
transfer protein treating atheros The compounds ha	Rodney, Burton 38 1 6562				
CAS INDEXING IS AVAILABLE FOR THIS PATENT.					
L21 ANSWER 16 OF 25 U ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S):	SPATFULL 1998:14840 USPATFULL Anthranilic acid derivatives Ozaki, Fumihiro, Ibaraki, Japan Ishibashi, Keiji, Ibaraki, Japan Ikuta, Hironori, Ibaraki, Japan Ishihara, Hiroki, Ibaraki, Japan Souda, Shigeru, Ibaraki, Japan Eisai Co., Ltd., Japan (non-U.S. corporation)				
	NUMBER KIND DATE				
PATENT INFORMATION: APPLICATION INFO.:	US 5716993 19980210 WO 9518097 19950706 US 1995-507476 19950914 (8) WO 1994-JP2262 19941227 19950916 PCT 371 date 19950916 PCT 102(e) date				
	NUMBER DATE				
PRIORITY INFORMATION: DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINER: LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT:	JP 1993-347092 19931227 JP 1994-299110 19941009 Utility Granted Owens, Amelia Nixon & Vanderhye 7 1 3902				

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides an anthranilic acid derivative AB having a cGMP-PDE inhibitory activity.

> An anthranilic acid derivative represented by the general formula (I) or a pharmacologically acceptable salt thereof: ##STR1## [wherein R.sup.1, R.sup.2, R.sup.3 and R.sup.4 represent the same or different from each other, a hydrogen atom, a halogen atom, a hydroxy group, an optionally halogenated lower alkyl group, an optionally halogenated lower alkoxy group, a nitro group, a hydroxyalkyl group, a cyano group or the like; R.sup.5 and R.sup.6 represent the same or different from each other, a hydrogen atom, a halogen atom, a hydroxy group, a cyano group, an optionally halogenated lower alkyl group, an optionally halogenated lower alkoxy group or the like;

W represents a group of the formula: --N.dbd. or --CH.dbd.; R.sup.7 and R.sup.8 represent the same or different from each other, a hydrogen atom, an optionally halogenated lower alkyl group or the like;

A represents a hydrogen atom, an optionally halogenated lower alkyl group or the like;

Y represents an oxygen atom or a sulfur atom; and

n is an integer of 0 to 6].

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 17 OF 25 USPATFULL

1998:9505 USPATFULL ACCESSION NUMBER:

TITLE:

Inhibitors of microsomal triglyceride transfer

protein and method

INVENTOR(S):

Biller, Scott A., Hopewell, NJ, United States Dickson, John K., Eastampton, NJ, United States Lawrence, R. Michael, Yardley, PA, United States Magnin, David R., Hamilton, NJ, United States Poss, Michael A., Lawrenceville, NJ, United

States

Robl, Jeffrey A., Newtown, PA, United States Sulsky, Richard B., Franklin Park, NJ, United

Tino, Joseph A., Lawrenceville, NJ, United States

Bristol-Myers Squibb Company, Princeton, NJ,

United States (U.S. corporation)

NUMBER KIND DATE ______ PATENT INFORMATION: US 5712279. 19980127

APPLICATION INFO.: RELATED APPLN. INFO.:

PATENT ASSIGNEE(S):

US 1996-548811 19960111 (8)

Continuation-in-part of Ser. No. US 1995-472067, filed on 6 Jun 1995 which is a

continuation-in-part of Ser. No. US 1995-391901,

filed on 21 Feb 1995, now abandoned

DOCUMENT TYPE: FILE SEGMENT:

Utility Granted

PRIMARY EXAMINER:

Shah, Mukund J.

308-4994 Shears Searcher :

ASSISTANT EXAMINER: LEGAL REPRESENTATIVE: Wong, King Lit

NUMBER OF CLAIMS:

Rodney, Burton 19

EXEMPLARY CLAIM:

1 2204

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compounds are provided which inhibit microsomal triglyceride transfer protein and thus are useful for lowering serum lipids and

treating atherosclerosis and related diseases.

The compounds have the structure ##STR1## wherein Z, X.sup.1,

X.sup.2, x and R.sup.5 are as defined herein.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 18 OF 25 USPATFULL

ACCESSION NUMBER:

97:16212 USPATFULL

TITLE:

Thioformamide derivatives

INVENTOR(S):

Kabasawa, Yasuhiro, Ibaraki, Japan Ozaki, Fumihiro, Ibaraki, Japan

Ishibashi, Keiji, Ibaraki, Japan Hasegawa, Takashi, Ibaraki, Japan Oinuma, Hitoshi, Ibaraki, Japan Ogawa, Toshiaki, Ibaraki, Japan Adachi, Hideyuki, Ibaraki, Japan Katoh, Hiroshi, Ibaraki, Japan Kodama, Kohtarou, Ibaraki, Japan Ohara, Hideto, Ibaraki, Japan Mori, Nobuyuki, Ibaraki, Japan Minami, Norio, Ibaraki, Japan

PATENT ASSIGNEE(S):

Eisai Co., Ltd., Japan (non-U.S. corporation)

NUMBER KIND DATE ______

PATENT INFORMATION: APPLICATION INFO .: US 5606061 19970225 US 1995-531335 19950920

RELATED APPLN. INFO.:

Division of Ser. No. US 1995-380589, filed on 30 Jan 1995, now patented, Pat. No. US 5498634 which is a division of Ser. No. US 1994-211701, filed on 26 Apr 1994, now patented, Pat. No. US 5444066

19920106

(8)

DATE NUMBER ______ JP 1991-264622 19911014

PRIORITY INFORMATION: DOCUMENT TYPE:

JP 1992-197 Utility Granted

FILE SEGMENT: PRIMARY EXAMINER: LEGAL REPRESENTATIVE:

Dentz, Bernard Nixon & Vanderhye

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 LINE COUNT: 2083

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides a thioformamide derivative AB represented by the following general formula (I) or a pharmacologically acceptable salt thereof, which is highly safe, easy to use, and useful as an excellent hypotensive or heart disease remedy: ##STR1## wherein Y represents ##STR2## or the like [wherein R.sup.7 represents benzyloxy or the like;

R.sup.11 and R.sup.12 each represent hydrogen, hydroxyl, benzoyloxy, benzyloxy, ##STR3## (wherein R.sup.14 and R.sup.15 each represent hydrogen, benzyl or the like) or the like);

Z represents --CH.sub.2 -- or the like; A represents imidazolyl or imidazopyridyl which may have one or two substituents, or the like; R.sup.1 and R.sup.2 each represent hydrogen, lower alkyl or the like; and R.sup.3 and R.sup.4 each represent hydrogen, lower alkyl or the like.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 19 OF 25 USPATFULL

ACCESSION NUMBER: 97:5853 USPATFULL

TITLE:

Nucleic acids encoding microsomal trigyceride

transfer protein

INVENTOR(S):

Wetterau, II, John R., Langhorne, PA, United

States

Sharp, Daru Y., Perrineville, NJ, United States Gregg, Richard E., Pennington, NJ, United States

Biller, Scott A., Ewing, NJ, United States

Dickson, John K., Mount Holly, NJ, United States Lawrence, R. Michael, Yardley, PA, United States Lawson, John E., Wallingford, CT, United States Holava, Henry M., Meriden, CT, United States Partyka, Richard A., Neshanic, NJ, United States

PATENT ASSIGNEE(S):

Bristol-Myers Squibb Company, Princeton, NJ,

Distor Chates (II C company) Firmon

United States (U.S. corporation)

	NUMBER	KIND	DATE	
US	5595872		19970121	
IIQ.	1993-117362		19930903	

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

US 1993-117362 19930903 (8) Continuation-in-part of Ser. No. US 1993-15449,

filed on 22 Feb 1993, now abandoned which is a continuation-in-part of Ser. No. US 1992-847503, filed on 6 Mar 1992, now abandoned

Utility

DOCUMENT TYPE: FILE SEGMENT:

Granted

PRIMARY EXAMINER:

Zitomer, Stephanie W. Gaul, Timothy J., Bogden, James M.

LEGAL REPRESENTATIVE:

14

NUMBER OF CLAIMS:

1

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

14 Drawing Figure(s); 8 Drawing Page(s)

LINE COUNT: 5232

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Nucleic acid sequences, particularly DNA sequences, coding for all or part of the high molecular weight subunit of microsomal triglyceride transfer protein, expression vectors containing the DNA sequences, host cells containing the expression vectors, and methods utilizing these materials. The invention also concerns polypeptide molecules comprising all or part of the high molecular weight subunit of microsomal triglyceride transfer protein, and methods for producing these polypeptide molecules. The invention additionally concerns novel methods for preventing, stabilizing or causing regression of atherosclerosis and therapeutic agents having such activity. The invention concerns further novel methods for lowering serum liquid levels and therapeutic

agents having such activity.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 20 OF 25 USPATFULL

96:21114 USPATFULL ACCESSION NUMBER:

Thioformamide derivatives TITLE:

Kabasawa, Yasuhiro, Ibaraki, Japan INVENTOR(S): Ozaki, Fumihiro, Ibaraki, Japan

Ishibashi, Keiji, Ibaraki, Japan Hasegawa, Takashi, Ibaraki, Japan Oinuma, Hitoshi, Ibaraki, Japan Ogawa, Toshiaki, Ibaraki, Japan Adachi, Hideyuki, Ibaraki, Japan Katoh, Hiroshi, Ibaraki, Japan Kodama, Kohtarou, Ibaraki, Japan Ohara, Hideto, Ibaraki, Japan

Mori, Nobuyuki, Ibaraki, Japan Minami, Norio, Ibaraki, Japan

Eisai Co., Ltd., Tokyo, Japan (non-U.S. PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE ______ US 5498634

19960312 PATENT INFORMATION: US 1995-380589

19950130 (8) APPLICATION INFO.:

Division of Ser. No. US 1994-211701, filed on 13 RELATED APPLN. INFO.:

Apr 1994, now patented, Pat. No. US 5444066

DATE NUMBER ______

JP 1991-264622 19911014 PRIORITY INFORMATION:

JP 1992-197 19920106 Utility DOCUMENT TYPE:

Granted FILE SEGMENT: PRIMARY EXAMINER: Dentz, Bernard

LEGAL REPRESENTATIVE: Nixon & Vanderhye NUMBER OF CLAIMS:

EXEMPLARY CLAIM: 2100 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides a thioformamide derivative represented by the following general formula (I) or a pharmacologically acceptable salt thereof, which is highly safe, easy to use, and useful as an excellent hypotensive or heart disease remedy: ##STR1## wherein Y represents ##STR2## or the like [wherein R.sup.7 represents benzyloxy or the like; R.sup.11 and R.sup.12 each represent hydrogen, hydroxyl, benzoyloxy, benzyloxy, ##STR3## (wherein R.sup.14 and R.sup.15 each represent hydrogen, benzyl or the like) or the like];

Z represents --CH.sub.2 -- or the like; A represents imidazolyl or imidazopyridyl which may have one or two substituents, or the like; R.sup.1 and R.sup.2 each represent hydrogen, lower alkyl or the like; and R.sup.3 and R.sup.4 each represent hydrogen, lower alkyl or the like.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Shears 308-4994 Searcher :

L21 ANSWER 21 OF 25 USPATFULL

ACCESSION NUMBER: 95:75975 USPATFULL

TITLE: Thioformamide derivatives having hypotensive

activity

INVENTOR(S): Kabasawa, Yasuhiro, Ibaraki, Japan

Ozaki, Fumihiro, Ibaraki, Japan Ishibashi, Keiji, Ibaraki, Japan Hasegawa, Takashi, Ibaraki, Japan Oinuma, Hitoshi, Ibaraki, Japan Ogawa, Toshiaki, Ibaraki, Japan Adachi, Hideyuki, Ibaraki, Japan Katoh, Hiroshi, Ibaraki, Japan Kodama, Kohtarou, Ibaraki, Japan Ohara, Hideto, Ibaraki, Japan Mori, Nobuyuki, Ibaraki, Japan

Minami, Norio, Ibaraki, Japan

PATENT ASSIGNEE(S): Eisai Co., Ltd., Tokyo, Japan (non-U.S.

corporation)

NUMBER DATE

PRIORITY INFORMATION: JP 1991-264622 19911014

JP 1992-4197 19920106

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

FILE SEGMENT: Granted
PRIMARY EXAMINER: Dentz, Bernard
LEGAL REPRESENTATIVE: Nixon & Vanderhye

NUMBER OF CLAIMS: 6
EXEMPLARY CLAIM: 1
LINE COUNT: 2109

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides a thioformamide derivative represented by the following general formula (I) or a pharmacologically acceptable salt thereof, which is highly safe, easy to use, and useful as an excellent hypotensive or heart disease remedy: ##STR1## wherein Y represents ##STR2## or the like [wherein R.sup.7 represents benzyloxy or the like; R.sup.11 and R.sup.12 each represent hydrogen, hydroxyl, benzoyloxy, benzyloxy, ##STR3## (wherein R.sup.14 and R.sup.15 each represent hydrogen, benzyl or the like);

Z represents --CH.sub.2 -- or the like; A represents imidazolyl or imidazopyridyl which may have one or two substituents, or the like; R.sup.1 and R.sup.2 each represent hydrogen, lower alkyl or the like; and R.sup.3 and R.sup.4 each represent hydrogen, lower alkyl or the like.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 22 OF 25 USPATFULL

ACCESSION NUMBER: 94:1438 USPATFULL

TITLE: Thioformamide derivative, process for its

preparation, pharmaceutical composition thereof

and treatment method

INVENTOR(S): Hart, Terance W., Brentwood, England

Vacher, Bernard Y. J., Dageham, England Walsh, Roger J. A., Rayleigh, England

PATENT ASSIGNEE(S): Rhone-Poulenc Sante, France (non-U.S.

corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 5276045 19940104
APPLICATION INFO.: US 1992-860599 19920330 (7)

RELATED APPLN. INFO.: Continuation of Ser. No. US 1990-538714, filed on

15 Jun 1990, now abandoned

NUMBER DATE

PRIORITY INFORMATION: GB 1989-13863 19890616

GB 1989-13864 19890616

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Shah, Mukund J.
ASSISTANT EXAMINER: Grumbling, Matthew V.
LEGAL REPRESENTATIVE: Morgan & Finnegan

NUMBER OF CLAIMS: 11
EXEMPLARY CLAIM: 1
LINE COUNT: 786

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A thioformamide derivative of the formula: ##STR1## wherein R represents alkyl, A represents optionally substituted pyrid-3-yl, isoquinolin-4-yl, tetrahydroquinolin-3-yl, quinolin-3-yl, pyridazin-4-yl, pyrimid-5-yl, thiazol-5-yl, thieno[2,3-b]pyridin-5yl, pyrazin-2-yl, indol-3-yl and thieno[3,2-b]pyridin-6-yl, or phenyl and Y represents a valency bond, methylene or ethylene, R.sup.2 represents hydrogen, optionally substituted alkyl, cycloalkyl, cycloalkylalkyl, aryl, aralkyl, aryloxyalkyl, aromatic heterocyclylalkyl or aromatic heterocyclyloxyalkyl group or a group ZC(.dbd.O) -- in which Z represents optionally substituted alkyl, aryl, or aromatic heterocyclic, n represents 0 or 1, and when n represents 0, R.sup.1 may represent a hydrogen atom, optionally substituted alkyl, cycloalkyl, cycloalkylalkyl, aryl, aralkyl, aryloxyalkyl, aromatic heterocyclylalkyl or aromatic heterocyclyloxyalkyl group or a group ZC(.dbd.0)-- or ZSO.sub.2 --, and when n represents 1, R.sup.1 represents optionally substituted alkyl, benzyl, phenethyl, 1-naphthylmethyl, 2-naphthylmethyl or pyrid-3-ylmethyl radical and pharmaceutically acceptable salts thereof possess pharmacological properties.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 23 OF 25 USPATFULL

ACCESSION NUMBER: 93:98381 USPATFULL

TITLE: Optionally substituted pyrido[2,3-d]pyridine-

2,4(1H,3H)-diones and pyrido[2,]pyrimidine-

2(1H, 3H) -ones

INVENTOR(S): Wilhelm, Robert S., Mountain View, CA, United

Chin, Ronnie L., Mountain View, CA, United States Devens, Bruce H., Palo Alto, CA, United States Alvarez, Robert, Menlo Park, CA, United States

PATENT ASSIGNEE(S): Syntex (U.S.A.) Inc., Palo Alto, CA, United

States (U.S. corporation)

NUMBER KIND DATE

US 5264437 19931123 PATENT INFORMATION: APPLICATION INFO.: US 1992-855179 · 19920320 (7)

Utility DOCUMENT TYPE: Granted FILE SEGMENT:

Page, Thurman K. PRIMARY EXAMINER: Venkat, Jyothsna ASSISTANT EXAMINER:

Wong, James J., Lowin, David A., Krubiner, Alan LEGAL REPRESENTATIVE:

Μ. 5.3 NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 3516 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to optionally substituted pyrido[2,3-d]pyrimidine-2,4(1H,3H)-diones or optionally substituted pyrido[2,3-d]pyrimidine-2(1H,3H)-ones, i.e., compounds of Formula I: ##STR1## wherein: Y is --CH.sub.2 -- or --C(O)--;

R.sup.1 is hydrogen or -- (CH.sub.2).sub.n --R.sup.7, wherein:

R.sup.7 is aryl or heteroaryl, and

n is 1 or 2,

provided that when Y is --C(0)--, R.sup.7 is heteroaryl; and

R.sup.2, R.sup.3, R.sup.4, R.sup.5 and R.sup.6 are hydrogen, or one is selected from lower alkyl, halo, carboxy, methoxycarbonyl, carbamoyl, methylcarbamoyl, di-methylcarbamoyl, methylcarbonyl, methylthio, methylsulfinyl, methylsulfonyl, hydroxymethyl, amino, trifluoromethyl, cyano or nitro; or

R.sup.2, R.sup.3, R.sup.4 and R.sup.5 are independently selected from hydrogen, lower alkyl, nitro, chloro, fluoro, methoxycarbonyl or methylcarbonyl, provided at least one is hydrogen, and R.sup.6 is hydrogen;

or a pharmaceutically acceptable ester, ether or salt thereof.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 24 OF 25 USPATFULL

ACCESSION NUMBER: 93:65400 USPATFULL

TITLE: Quinazoline-3-alkanoic acid derivatives, their

salts and their preparation processes Fujimori, Shizuyoshi, Marubayashi, Japan

INVENTOR(S): Ohnota, Michiro, Nogi, Japan

Hirata, Yoshihiro, Omiya, Japan

Murakami, Koji, Nogi, Japan

PATENT ASSIGNEE(S): Kyorin Pharmaceutical Co., Ltd., Tokyo, Japan

(non-U.S. corporation)

KIND DATE NUMBER ______ US 5234928 19930810 PATENT INFORMATION: WO 9109024 19910627 APPLICATION INFO.: US 1991-721610 19910717 (7) WO 1990-JP1600 19901210 19910717 PCT 371 date 19910717 PCT 102(e) date

NUMBER DATE

PRIORITY INFORMATION: JP 1989-321097 19891211

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NUMBER OF CLAIMS: 3
EXEMPLARY CLAIM: 1
LINE COUNT: 1070

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to quinazoline-3-alkanoic acid derivatives having both an inhibitory effect on platelet aggregation and a hindering effect on aldose reductase together, represented by a general formula [I] ##STR1## [wherein R is hydrogen or a protecting group for carboxyl group, R.sup.1 is a lower alkyl group, alkenyl group, alkinyl group, lower alkoxy group, lower alkylthio group, halogen, phenyl group (this phenyl group may be substituted by one to three of lower alkyls, lower alkoxys, halogens, trifluoromethyls, carboxyethylenes or ethoxycarbonylethylenes, naphthyl group, heterocycle (this heterocycle may be substituted by one to three of lower alkyls), cycloalkyl group or benzoyl group (this benzoyl group may be substituted by lower alkyl or halogen), R.sup.2 and R.sup.3 are identically or differently hydrogens, halogens, lower alkyl groups, lower alkoxy groups, aralkyl groups which may be substituted, nitro groups, imidazolyl groups, imidazolylmethyl groups or ##STR2## (R.sup.4 and R.sup.5 indicate identically or differently hydrogens or lower alkyl groups, or connected with each other to make five- or six-membered heterocycles which may contain other hetero atom, X is carbonyl, thiocarbonyl or methylene group (this methylene group may be substituted by lower alkyl group), A is lower alkylene or lower alkenylene, and n indicates an integer of 1 to 3],

their salts, their preparation processes and medicinal drugs containing them.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L21 ANSWER 25 OF 25 USPATFULL

ACCESSION NUMBER: 86:29854 USPATFULL

TITLE: Acridinecarboxamide compounds

INVENTOR(S): Atwell, Graham J., Auckland, New Zealand

Baguley, Bruce C., Auckland, New Zealand Denny, William A., Auckland, New Zealand Rewcastle, Gordon W., Auckland, New Zealand Development Finance Corporation of New Zealand,

PATENT ASSIGNEE(S): Development Finance Corporation of New Zealand (non-U.S. corporation)

NUMBER DATE

PRIORITY INFORMATION: NZ 1982-201084 19820625

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FILE SEGMENT: Granted
PRIMARY EXAMINER: Daus, Do

PRIMARY EXAMINER: Daus, Donald G.
ASSISTANT EXAMINER: Rivers, Diana G.
LEGAL REPRESENTATIVE: Pennie & Edmonds

NUMBER OF CLAIMS: 7
EXEMPLARY CLAIM: 1
LINE COUNT: 1063

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB 4-Carboxamidoacridine compounds represented by the general formula (I), ##STR1## where R.sub.1 represents H, CH.sub.3 or NHR.sub.3, where R.sub.3 is H, COCH.sub.3, SO.sub.2 CH.sub.3, COPh, SO.sub.2 Ph or lower alkyl optionally substituted with hydroxyl and/or amino functions;

R.sub.2 represents H or up to two of the groups CH.sub.3, OCH.sub.3, halogen, CF.sub.3, NO.sub.2, NH.sub.2, NHCOCH.sub.3, and NHCOOCH.sub.3 placed at positions 1-3 or 5-8;

Y represents C(NH)NH.sub.2, NHC(NH)NH.sub.2, or NR.sub.4 R.sub.5, where each of R.sub.4 and R.sub.5 is H or lower alkyl optionally substituted with hydroxyl and/or amino functions; and

x is from 2 to 6,

and the acid addition salts thereof, possess antibacterial and antitumor properties.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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